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1 Biological Sciences – Microbiology

2 **Interplay between tolerance mechanisms to copper and acid stress in *Escherichia***

3 ***coli***

4

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23 biosynthesis

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25

1 **ABSTRACT (217 words)**

2 Copper (Cu) is a key antibacterial component of the host innate immune
3 system and almost all bacterial species possess systems that defend against the toxic
4 effects of excess Cu. The Cu tolerance system in Gram-negative bacteria comprises
5 minimally of a Cu sensor (CueR) and a Cu export pump (CopA). The *cueR* and *copA*
6 genes are encoded on the chromosome typically as a divergent but contiguous operon.
7 In *E. coli*, *cueR* and *copA* are separated by two additional genes, *ybaS* and *ybaT*,
8 which confer glutamine (Gln)-dependent acid tolerance and contribute to the
9 glutamate (Glu)-dependent acid resistance system in this organism. Here we show that
10 Cu strongly inhibits growth of a $\Delta copA$ mutant strain in acidic cultures. We further
11 demonstrate that Cu stress impairs the pathway for Glu biosynthesis *via* glutamate
12 synthase (GltBD or GOGAT), leading to decreased intracellular levels of Glu.
13 Addition of exogenous Glu rescues the $\Delta copA$ mutant from Cu stress in acidic
14 conditions. Gln is also protective but this relies on the activities of YbaS and YbaT.
15 Notably, expression of both enzymes is upregulated during Cu stress. These results
16 demonstrate a link between Cu stress, acid stress, and Glu/Gln metabolism, establish a
17 role for YbaS and YbaT in Cu tolerance, and suggest that subtle changes in core
18 metabolic pathways may contribute to overcoming host-imposed copper toxicity.

19

1 **SIGNIFICANCE STATEMENT (101 words)**

2 Copper is an essential trace metal nutrient in health and is increasingly
3 recognized for its role in the control of infection. The pathogen *Escherichia coli*
4 encounters host niches with mild to high acidity and elevated copper levels. Our study
5 shows that this bacterium can alter its metabolism and harness the amino acid
6 glutamine to suppress the effects of acid stress and copper toxicity. Given the
7 abundance of glutamine in systemic circulation and its importance in the host immune
8 system, our work provides a new insight into the ways in which bacterial pathogens
9 can adapt and survive host-imposed antibacterial strategies.

1 \body

2 INTRODUCTION

3 The efflux of excess transition metal ions such as copper (Cu) is an important
4 feature of bacterial physiology, particularly during the interactions between a bacterial
5 pathogen and its host. Several lines of evidence have established a role for Cu as a
6 host-derived antibacterial agent that contributes to nutritional immunity (1). In turn,
7 the ability to export Cu from the bacterial cytoplasm is now recognised as a key
8 determinant of bacterial virulence (2-5). At the biochemical level, the mechanisms of
9 bacterial Cu export are well understood and, in Gram-negative species, are
10 exemplified by the Cue/Cop regulon in *Escherichia coli* (6). This system consists of a
11 Cu(I)-sensing transcriptional regulator (CueR), which controls expression of a
12 transmembrane efflux pump (CopA) that exports Cu(I) from the cytoplasm to the
13 periplasm and a periplasmic cuprous oxidase (CueO) that converts Cu(I) to the less
14 toxic Cu(II) form.

15 In the *E. coli* chromosome, *copA* and *cueR* are separated by an operon
16 annotated as *ybaST*, which is encoded in the same orientation as *cueR* and is divergent
17 from *copA* (Fig. 1A). *ybaS* encodes a glutaminase that catalyses the hydrolysis of
18 *L*-glutamine (Gln) to generate *L*-glutamate (Glu) and ammonia (Fig. 1B). The
19 glutaminase activity of YbaS confers Gln-dependent acid tolerance and contributes to
20 the Glu-dependent system for acid resistance (AR) (7, 8). In this AR mechanism, Glu
21 is converted to γ -aminobutyric acid (GABA) by two glutamate decarboxylases
22 (GadA, GadB). This process consumes a proton and raises the cytoplasmic pH (9).
23 The intracellular Glu pool can be replenished if an extracellular supply is present and
24 this import occurs *via* the permease GadC (Fig. 1B). *ybaT* encodes an amino acid

1 permease that may also contribute to Glu-dependent AR by supplying Gln to YbaS for
2 hydrolysis (Fig. 1B) but its substrate specificity remains to be established .

3 Here we examine the significance of the synteny of Cu tolerance and acid
4 tolerance genes in relation to *E. coli* physiology. Using a $\Delta copA$ deletion mutant
5 strain, we show a link between Cu stress, acid stress, and Glu/Gln metabolism, and
6 establish a role for YbaS and YbaT in Cu tolerance. Our results suggest that subtle
7 changes in bacterial metabolism may contribute to overcoming host-imposed copper
8 toxicity during nutritional immunity.

10 RESULTS

11 **Organisation of *copA* and *cueR* in *E. coli*.** The nucleotide sequence of the
12 *copA-ybaST-cueR* locus from *E. coli* was used to query all complete bacterial genome
13 sequences in the NCBI database (see Supporting Methods in the SI Appendix). The
14 search results indicated that the synteny of *copA*, *ybaST*, and *cueR* is unique to
15 *Escherichia* and *Shigella* genera (Fig. S1). In the case of *E. coli*, this gene cluster is
16 part of the core genome (10) and is found in 208/209 complete genomes (>95%
17 sequence conservation), which include strains representative of environmental,
18 commensal, and all pathogenic types.

19 The *ybaST* insertion is absent from other *Enterobacteriaceae* such as
20 *Salmonella* and *Klebsiella*. In these organisms, *copA* is divergent from but contiguous
21 with *cueR* (Fig. S2), which is the canonical arrangement for a *merR*-like operon. One
22 exception was *Serratia marcescens*, in which *cueR* and *copA* are separated by a
23 cluster of genes encoding for the biosynthesis of the antibiotic prodigiosin (*pigA-O*).
24 Expression of *pig* genes is repressed by Cu but the physiological relevance of this
25 observation is unclear (11). In agreement with a previous report, *ybaS* is also present

1 in nine additional genera from the *Enterobacteriaceae* family, including *Edwardsiella*
2 and *Yersinia*. However, consistent with its established function in acid tolerance, *ybaS*
3 in these genomes is frequently encoded adjacent to *gadA/B* or *gadC* genes for AR
4 (Fig. S3).

5

6 **Cu stress in a $\Delta copA$ mutant is enhanced during growth in acidic**

7 **conditions.** To determine if there was a link between Cu and acid stress, we examined
8 the inhibitory effects of added Cu on the growth of *E. coli* in minimal medium
9 buffered at pH 5 and pH 7. Addition of up to 1.0 μ M Cu did not impact growth of the
10 wild type (WT) strain at either pH but it inhibited growth of the $\Delta copA$ mutant (Fig.
11 2). Notably, the amount of Cu required to completely suppress growth at pH 5 (0.1
12 μ M, Fig. 2A) was less than the amount required at pH 7 (1.0 μ M, Fig. 2B), suggesting
13 that Cu stress in the $\Delta copA$ mutant was enhanced during growth in acidic conditions.
14 The Cu-tolerant phenotype was restored upon expression of *copA* via plasmid-
15 mediated complementation (Fig. 2). Identical results were obtained using the $\Delta copA$
16 mutants of other pathogenic and nonpathogenic strains of *E. coli* (Fig. S4), indicating
17 that the interplay between Cu stress and acid stress is a conserved feature of *E. coli*
18 physiology.

19 Addition of Cu to 0.05 μ M was sufficient to affect growth at pH 5 and not at
20 pH 7 (cf. Fig. S4, Fig. 2). However, this treatment led to a comparable rise in total
21 intracellular Cu levels in mid-exponential $\Delta copA$ cells during growth at both pH
22 values as determined by ICP MS (Fig. S5). Hence, the increase in Cu stress at pH 5
23 did not correlate with an increase in the amounts of trapped Cu. Nevertheless, Cu may
24 become more bioavailable during growth at pH 5 due to protonation of thiols and
25 amines, leading to a decrease in the Cu buffering capacity of the extracellular medium

1 or intracellular milieu. To report for bioavailable Cu, we used a *lacZ* transcriptional
2 reporter fused to the *copA* promoter (P_{copA} -*lacZ*) (12). The latter is activated by CueR,
3 the primary Cu sensor in *E. coli* (Fig. 1B). In agreement with a previous observation
4 (13), background P_{copA} -*lacZ* activity in mid-exponential $\Delta copA$ cells was higher than
5 in WT and it did not increase further in response to added Cu (Fig. S6A), suggesting
6 that this mutant trapped trace amounts of Cu in its cytoplasm. Nevertheless, P_{copA} -*lacZ*
7 activities in the $\Delta copA$ mutant were comparable irrespective of the pH of the culture
8 medium (Fig. S6A). Similarly, the response of the P_{copA} -*lacZ* reporter to Cu was
9 comparable in WT cultures grown at pH 5 and pH 7 (Fig. S6B). Thus, the present
10 evidence did not support an increased cellular level or bioavailability of Cu at pH 5
11 under the experimental conditions employed here.

13 Exogenous Glu and Gln protect the $\Delta copA$ mutant from Cu stress at pH 5.

14 Given the established role of *ybaS* in Gln-dependent acid tolerance, its demonstrated
15 ability to contribute to Glu-dependent AR, and its synteny with *copA* and *cueR* (Fig.
16 1), we hypothesised that Cu stress during growth in acidic conditions is linked to Glu
17 and/or Gln utilization. Indeed, addition of exogenous Glu and Gln (0.5 mM total)
18 rescued the growth of the $\Delta copA$ mutant in Cu-supplemented medium at pH 5 (Fig.
19 3A) and pH 7 (Fig. 3B). This protection was also observed for other pathogenic and
20 nonpathogenic strains of *E. coli* (Fig. S7). Compared with Gln, protection by Glu
21 extended to higher Cu concentrations (Fig. S8).

22 Asp and Asn were also protective at pH 5 (Fig. S9A). In contrast, Arg, which
23 is responsible for an alternative acid resistance system in *E. coli* (9), was only
24 partially protective at pH 5 (Fig. S9B). Cys and the Cys-containing tripeptide
25 glutathione were strongly protective (Fig. S9C), likely because these thiol-containing

1 molecules are high-affinity chelators of Cu. We also tested branched-chain amino
2 acids (BCAAs, *ie.* Ile, Leu, and Val), which are known to rescue Cu-sensitive mutants
3 of *E. coli* from Cu stress at pH 7 (13, 14). These amino acids rescued growth of the
4 $\Delta copA$ mutant strongly at pH 7 (Fig. 3B) but only weakly at pH 5 (Fig. 3A),
5 especially when compared with Glu and/or Gln. These results indicated that Cu stress
6 at pH 5 led to a high requirement for exogenous Glu and/or Gln but not BCAAs.
7 Nevertheless, addition of Glu and/or Gln did not restore Cu tolerance of the $\Delta copA$
8 mutant to WT levels. This observation and the varying protective effects of each
9 amino acid are discussed below.

10

11 **Cu stress leads to depletion in intracellular Glu concentrations.** Glutamate
12 is a key metabolite in *E. coli*, accounting for nearly 40% of all intracellular
13 metabolites during exponential growth (15). The observed auxotrophy for Glu during
14 Cu stress (Fig. 3) indicated that the internal pool of this amino acid was depleted. To
15 test this hypothesis, *E. coli* was cultured with or without added Cu to the
16 mid-logarithmic phase and the concentrations of free Glu in harvested cells were
17 measured. As anticipated, growth in Cu-supplemented medium diminished the Glu
18 pools in the $\Delta copA$ mutant (Fig. 4A). This effect was observed at both pH 5 (Fig.
19 4A(ii)) and pH 7 (Fig. 4A(i)) but less Cu was required at pH 5, consistent with our
20 earlier finding that less Cu was required to inhibit the $\Delta copA$ mutant at pH 5 (Fig. 2).
21 Intracellular Glu levels in WT cells remained unaffected under these conditions (Fig.
22 4A).

23 Our measurements further revealed that basal Glu levels in mid-exponential
24 $\Delta copA$ cells cultured at pH 5 without any added Cu (~9 nmol Glu/mg protein, Fig.
25 4A(ii)) were lower than in cells cultured at pH 7 (~44 nmol Glu/mg protein, Fig.

1 4A(i)). This Glu-starved phenotype was likely associated with the consumption of this
2 amino acid to maintain the internal pH during growth in mild acid, which could occur
3 *via* the GadA/B decarboxylases (*cf.* Fig. 1B) (16, 17). Hence, the observed importance
4 of exogenous Glu but not BCAAs or other amino acids during Cu stress at pH 5 (*cf.*
5 Fig. 3A) may reflect the unique role for Glu for growth in acidic conditions.

6

7 **Excess Cu inhibits Glu biosynthesis *via* glutamate synthase (GOGAT).**

8 Glutamate in *E. coli* is synthesized from α -ketoglutarate (α -KG), an intermediate in
9 the TCA cycle, *via* two pathways (Fig. 4B). Glutamate dehydrogenase (GDH or
10 GdhA) generates Glu from the reductive amination of α -KG. In the alternative
11 pathway, condensation of Glu with ammonia by glutamine synthetase (GS or GlnA)
12 yields Gln. Subsequent reductive transamination of Gln with α -KG by glutamate
13 synthase (GOGAT or GltBD) generates two Glu molecules (a net gain of one).
14 Together, GDH, GS, and GOGAT constitute the central pathway for nitrogen
15 assimilation in *E. coli*. The GDH route is thought to be most efficient when ammonia
16 is abundant, while GOGAT is important during ammonia limitation (18).

17 Addition of the Glu precursor α -KG failed to protect the $\Delta copA$ mutant from
18 Cu stress at pH 5 and pH 7 (Fig. 3), implying that either GDH or GOGAT, or both,
19 was inactive. Growth at pH 5 in the presence of added Cu did not reduce *gdhA*
20 transcription (Fig. 4C(i)) or GDH activity (Fig. 4D(i)) in mid-exponential $\Delta copA$
21 cells. At pH 7, there was a decrease in *gdhA* transcription (Figures 4C(i)) but there
22 was no loss in GDH activity (Fig. 4D(i)). By contrast, growth in Cu-supplemented
23 medium at pH 5 reduced the activity of GOGAT in mid-exponential $\Delta copA$ cells by
24 ca. 50% (Fig. 4D(ii)). GOGAT activity was also lost during growth in Cu-
25 supplemented medium at pH 7 but the amount of Cu required to achieve this effect

1 was again higher than at pH 5 (Fig. 4D(ii)). There was no change in the levels of *gltB*
2 transcription in these Cu-treated cultures (Fig. 4C(ii)), suggesting that Cu exerted an
3 inhibitory effect at the protein level. Both GOGAT and GDH remained active in WT
4 cells during growth at either pH, with or without added Cu (Fig. S10).

5 Cu-stressed $\Delta copA$ cultures displayed additional phenotypes consistent with
6 GOGAT deficiency. There was a small but reproducible increase in GDH activity in
7 $\Delta copA$ cells grown in Cu-rich medium at pH 5 (Fig. 4D(i)). Upregulation of GDH has
8 been shown to compensate for the decrease in Glu biosynthesis in $\Delta gltD$ and $\Delta gltB$
9 mutant strains of *E. coli* and *Salmonella*, respectively (19). Decreasing the amount of
10 ammonia in the culture medium further exacerbated Cu stress in $\Delta copA$ cells at both
11 pH 7 and pH 5 (Fig. S11), consistent with GDH becoming more inefficient and
12 increased reliance on GOGAT for making Glu (18) (*cf.* Fig. 4B).

13
14 **Mechanism of GOGAT inhibition.** GOGAT contains two [4Fe-4S] clusters
15 in the active site. The decrease in GOGAT activity during Cu stress at pH 5 and pH 7
16 corresponded with reductions in the activities of two additional [4Fe-4S]-containing
17 enzymes, namely NADH dehydrogenase I (NUO) and succinate dehydrogenase
18 (SDH) (Fig. S12). These results suggested that Cu stress in our experimental
19 conditions interfered with the synthesis, maturation, and incorporation of Fe-S
20 clusters into enzymes *via* the Isc pathway as proposed previously (13, 14, 20). Yet,
21 contrary to these prior reports, the amounts of Cu used in these experiments were not
22 sufficient to induce expression of *sufA* and *sufB*, two genes that encode components of
23 the Suf pathway for Fe-S cluster repair (Fig. S13A-B, discussed below).

24 Mature, solvent-accessible [4Fe-4S] clusters are also known to be destroyed
25 by excess Cu (13, 21). To determine if GOGAT was poisoned directly by Cu, the

1 *ΔcopA* mutant was cultured without any added Cu and, upon reaching mid-
2 exponential phase, was exposed to added Cu (0–10 μM) for 30 min. Consistent with
3 our hypothesis, this treatment had no effect on GDH activity but it reduced GOGAT
4 activity in a dose-dependent manner (Fig. S14), suggesting that at least one of its
5 [4Fe-4S] clusters is solvent-exposed. As a control, we confirmed that the activity of
6 NUO remained unaffected (Fig. S14), likely because its [4Fe-4S] clusters are
7 protected by the protein scaffold. However, the doses of Cu that poisoned GOGAT
8 (~10 μM) were higher than doses that suppressed growth (< 1 μM, cf. Fig. 2, Fig. S4).
9 Hence, we propose that the loss of GOGAT during Cu stress was associated primarily
10 with the loss of [4Fe-4S] cluster biosynthesis (also see Supporting Discussion 1).

11
12 **An implication for the loss of GOGAT at pH 5.** The varying protective
13 effects of amino acids during Cu stress as described earlier can be reconciled with an
14 impaired GOGAT activity. Exogenous Glu was protective (Fig. 3) likely because it
15 bypassed GOGAT and supplied cells with the final reaction product (*cf.* Fig. 4B).
16 Similarly, Asp and Asn were protective (Fig. S9A) because these amino acids could
17 act as substrates for an alternative route to Glu. The three enzymes involved in this
18 pathway, aspartate aminotransferase (AspC) and the two asparaginases (AnsA and
19 AnsB, Fig. S9A), are presumably Cu-tolerant because they do not require Fe-S
20 clusters or other transition metal ions as cofactor.

21 As noted earlier, exogenous Glu did not restore Cu tolerance of the *ΔcopA*
22 mutant to WT levels, likely because multiple other Fe-S clusters were also inactivated
23 (*eg.* NUO and SDH). While addition of Glu would alleviate the demand on GOGAT
24 and upstream Fe-S enzymes that ultimately feed into the Glu pool (*eg.* aconitase), it
25 would not relieve the block on other defective enzymes such as the Fe-S dehydratases

1 in the pathway for BCAA biosynthesis. Similarly, addition of BCAAs only partially
2 rescued the $\Delta copA$ mutant from Cu stress (Fig. 3), likely because GOGAT and other
3 Fe-S enzymes outside the BCAA biosynthesis pathway remained inactive.

4 It is important to highlight that at pH 5, the protective effect of BCAAs was
5 diminished relative to Glu (Fig. 3A). This observation was consistent with the
6 established requirement for Glu, but not BCAAs, for acid tolerance (*cf.* Fig. 1B). The
7 K_m values for the two Glu decarboxylases in *E. coli* have been reported to range
8 between 1 to 15 mM (22-24). Our estimate of the intracellular Glu pool at pH 5 was
9 within this range (~9 nmol/mg protein or ~3 mM). Hence, we anticipate that the
10 inability to synthesise Glu *via* GOGAT (and the corresponding decrease in the Glu
11 pool) would impact the efficiency of acid tolerance under our experimental
12 conditions.

13

14 **Cu stress induces expression of YbaS and YbaT.** Exogenous Gln was also
15 protective, albeit to a lesser extent compared with Glu (Fig. S8). This observation was
16 counterintuitive given that Gln is a co-substrate for GOGAT (Fig. 4B). However, like
17 Asp and Asn, Gln can be converted to Glu independently from GOGAT by two
18 separate glutaminases, namely YbaS (glutaminase A) and YneH (glutaminase B) (Fig.
19 4B). Although Cu stress did not affect YneH activity in mid-exponential $\Delta copA$ cells
20 (Fig. 4D(iii)), it increased the activity of YbaS (Fig. 4D(iv)). This increase was
21 observed during growth at pH 5 and pH 7, although again the amount of Cu required
22 to achieve this effect was less at pH 5 (Fig. 4D(iv)). The upregulation in YbaS activity
23 correlated with an increase in the levels of *ybaS* (Fig. 4C(iv)) and *ybaT* (Fig. S13C)
24 expression. In contrast, expression of *yneH* was not altered in response to Cu (Fig.
25 4C(iii)).

1 In the simplest model, induction of *ybaST* in response to Cu stress would
2 occur in a CueR-dependent manner (*cf.* Fig. 1). To test this hypothesis, we constructed
3 a $\Delta copA\Delta cueR$ double mutant. This mutant was reproducibly more Cu-sensitive than
4 was the $\Delta copA$ parent strain at pH 5 (Fig. S15 and Supporting Discussion 2).
5 Nevertheless, Cu treatment also increased the amounts of *ybaST* transcripts in the
6 $\Delta copA\Delta cueR$ mutant when compared with the untreated control (Fig. S16), suggesting
7 that CueR did not directly regulate expression of *ybaST*, at least under our
8 experimental conditions.

9 Although the *ybaS* and *ybaT* genes for acid tolerance were induced during Cu
10 stress, the *copA* gene for Cu tolerance did not appear to be upregulated during acid
11 stress. Basal expression levels of *copA* (Fig. S17) and activities of the P_{copA} -*lacZ*
12 fusion (Fig. S6) in mid-exponential WT cells were comparable regardless of the
13 growth pH. Thus, the evidence described in this work collectively point to an indirect
14 and potentially complex relationship between Cu tolerance and acid tolerance
15 systems.

16

17 **The protective effect of Gln is suppressed in a $\Delta ybaST$ mutant.** Our results
18 led us to propose that YbaS and YbaT act as a compensatory pathway that offsets the
19 loss in GOGAT and protects the intracellular Glu pool *via* hydrolysis of Gln and
20 regeneration of Glu (Figs. 1B, 4B). The glutaminase activity of YbaS also produces
21 ammonia, which contributes to overall acid tolerance (Fig. 1B) (7). This model
22 predicts that the protective effect of exogenous Gln would be suppressed if YbaS and
23 YbaT were inactive. To test this proposal, we mutated the entire *ybaST* operon in the
24 $\Delta copA$ genetic background. The resulting $\Delta copA\Delta ybaST$ mutant strain was confirmed
25 to display no measurable YbaS activity (Fig. S18).

When cultured in the absence of added Gln, the $\Delta copA\Delta ybaST$ mutant displayed a Cu-sensitive phenotype that was comparable to the $\Delta copA$ parent strain (Fig. S19). However, when the growth medium at pH 5 was supplemented with Gln (0.5 mM), the $\Delta copA\Delta ybaST$ double mutant was demonstrably more Cu-sensitive than was the $\Delta copA$ parent strain, as evidenced by a prolonged lag phase in the presence of Cu (Fig. 5A). This observation indicated that the absence of functional *ybaS* and *ybaT* genes led to an increased sensitivity to Cu in the presence of Gln, or alternatively, a diminished protective effect of Gln during Cu stress. This phenotype was observed regardless of ammonia availability (Fig. S20). Expression of the *ybaST* operon on a plasmid fully restored YbaS activity (Fig. S18) and, subsequently, the protective effect of Gln (Fig. 5A). By contrast, deletion of *ybaST* had only a minor impact on Gln protection at pH 7, suggesting that this operon plays a less important role at pH 7 (Fig. 5B). These results confirmed that the protective effect of Gln against Cu stress during growth in acidic conditions required *ybaS* and *ybaT*, and they established a link between Cu stress, acid tolerance, and Gln utilisation.

DISCUSSION

The antibacterial activity of Cu is an important component of the mammalian innate immune defense (1). In response, bacterial pathogens mount a survival strategy that relies on the efflux of excess Cu ions from their cytoplasm. The *copA* genes of clinically significant pathogens, including *Mycobacterium tuberculosis* (3), *Streptococcus pneumoniae* (2), and *Klebsiella pneumoniae* (4), have been identified as a virulence factor in animal models of infection. We have previously described the synergistic action of Cu ions with other antibacterial agents that may be derived from

1 the host, such as nitric oxide (25) and hydrogen peroxide (26). Here we provide
2 evidence that Cu ions and acid are also strong co-stressors.

3 Cu is a highly competitive metal that outcompetes weaker binding metals from
4 sites in metalloproteins, leading to Cu intoxication in cells. Several proteins that
5 contain Fe (particularly Fe-S clusters) (13, 14, 21, 26, 27), Zn (28), and Mn (29) have
6 now been identified as targets of Cu poisoning. Since metalloproteins account for
7 nearly half of all enzymes in cells (30), precisely which enzymes are mismetallated by
8 Cu and the ensuing changes in bacterial physiology may vary, depending on the
9 specific organism and experimental conditions. The latter do not always approximate
10 the natural environment of the organism under investigation. In the case of *E. coli*, the
11 unprecedented synteny of Cu tolerance genes with Gln-dependent acid tolerance
12 genes in its chromosome may provide an insight. Importantly, this genetic
13 arrangement is conserved in *E. coli*, implying strong selection pressure.

14 *E. coli* resides primarily in the lower intestines of mammals. During its
15 interaction with the animal host, this bacterium experiences mild and extreme
16 fluctuations in external pH in the stomach, intestinal lumen, genitourinary tract, and
17 phagolysosomes of epithelial and innate immune cells. Recent evidence suggests that
18 *E. coli* also encounters elevated levels of Cu in at least some of these sites. Survival of
19 a $\Delta copA$ mutant within murine macrophages was impaired when compared with the
20 WT (5). In uropathogenic *E. coli*, *copA*, as well as *cueO*, *cusC*, and *cusF* genes for Cu
21 tolerance were highly expressed during human urinary tract infection, and this
22 observation correlated with increased Cu concentrations in the urine of infected
23 patients (31).

24 The combination of Cu and low pH poses a unique challenge to *E. coli*
25 metabolism. Our *in vitro* work showed that excess Cu ions in the *E. coli* cytoplasm

1 may impair acid tolerance by disrupting Glu biosynthesis *via* GOGAT. However, our
2 data also suggested that *E. coli* may use alternative enzymes, namely YbaS and YbaT,
3 to overcome this block in Glu synthesis if exogenous Gln is supplied. Intriguingly,
4 Gln is the most abundant amino acid in systemic circulation. Approximately 600 μ M
5 of free Gln is present in human blood plasma and nearly a third of this supply is
6 turned over by gastrointestinal mucosa epithelial cells (32, 33). Gln is also
7 indispensable for the proliferation and antimicrobial activity of innate immune cells
8 (34). It is plausible that to survive *in vivo*, *E. coli* can alter its metabolism to access
9 host Gln stores using the mechanisms identified in this work. Notably, both *ybaS* and
10 *ybaT* were also upregulated along with *copA* during extraintestinal urinary tract
11 infection in humans (31).

12 Other enteric bacteria, for example *Salmonella* and *Klebsiella*, share common
13 colonisation routes and niches with *E. coli*, and they are presumably also exposed to
14 the combination of acid and Cu stress. Like *E. coli*, both organisms use GOGAT to
15 synthesise Glu. However, neither relies on Glu for acid tolerance (35) and thus
16 poisoning of the Glu biosynthesis pathway would not have the same impact on their
17 survival. It is worth noting that in these organisms, *copA* and *cueR* are contiguous, and
18 homologues for YbaS and YbaT are absent (Fig. S2). The key targets of Cu stress in
19 these organisms remain to be identified but for *Klebsiella pneumoniae*, [4Fe-4S]
20 dehydratases in the pathway for branched-chain amino acid biosynthesis are major
21 candidates, at least during colonisation of the lung (4).

22 How is *ybaST* upregulated by Cu? The intergenic region between *copA* and
23 *ybaST* contains three regulatory sites (Fig. 1A): CueR, the primary Cu sensor in *E.*
24 *coli* and regulator of *copA*; CpxR, which controls the global response to envelope
25 stress; and GadX/W, which governs transcription of acid resistance genes including

1 *gadA/B* and *gadC*. MerR-like regulators regulate transcription of divergent sequences
2 in both directions (36) but CueR does not appear to also control *ybaST* (Fig. S16).
3 Meanwhile, the putative role of CpxR in *ybaST* transcription has not been
4 experimentally tested (37). In contrast, regulation of *ybaST* by GadX and GadW is
5 well established, and this occurs in a RpoS-dependent manner (38-40). Notably, RpoS
6 and CpxR also contribute to the cellular adaptation to excess Cu (41, 42). Finally, our
7 results link Cu stress to central nitrogen metabolism and thus additional control by the
8 nitrogen response regulators, such as NtrB and NtrC, may be involved (18). These
9 regulators sense intracellular Gln/Glu/ α -KG ratios, which may shift during conditions
10 of Cu stress (*cf.* Fig. 4A). Ntr regulation is further coupled with RpoS *via* the stringent
11 response and the (p)ppGpp signaling alarmone (18, 43). It is likely that the *ybaST*
12 operon is subject to a complex network of regulatory controls not unlike those
13 identified for the *gadA/BC* genes for Glu-dependent AR. Further studies are required
14 to elucidate how these hierarchies of regulation are coordinated.

15 Finally, we noted that Cu stress in our experimental conditions was sufficient
16 to induce *ybaS* and *ybaT* but not the *suf* pathway for the repair of damaged iron-sulfur
17 clusters (Fig. S13). Hence, we propose that upregulation of *ybaS* and *ybaT* represents
18 subtle metabolic changes that may occur in the graded response to low physiological
19 levels of Cu encountered in the host.

20

21 **METHODS**

22 **Strains and culture conditions.** *E. coli* EC958, UTI89, and MG1655 were
23 used in this study as indicated. All strains were propagated from frozen glycerol
24 stocks on antibiotic-free LB agar or, where specified, in liquid M9 medium. Strains
25 carrying pSU2718 plasmids were propagated in the presence of chloramphenicol (30

1 $\mu\text{g/mL}$). Liquid cultures were prepared in modified M9 medium (3 g/L KH_2PO_4 , 0.5
2 g/L NaCl, 1 mM MgSO_4 , 0.1 mM CaCl_2 , 16.5 $\mu\text{g/mL}$ thiamine, 25 mg/mL
3 nicotinamide) using glucose (1 g/L) as carbon source and ammonium chloride (1 g/L)
4 as nitrogen source. The medium was buffered at pH 5 with Na-MES (50 mM) or at
5 pH 7 with Na-MOPS (50 mM). These pH buffers do not form a stable complex with
6 Cu^{2+} ions (44). The medium was used without any metal purification step. Cultures
7 were inoculated to an initial OD_{600} of 0.01 and grown at 37 °C with shaking at 200
8 rpm.

9 **Construction of mutants.** Deletion mutants of *copA*, *cueR*, and the *ybaST*
10 operon were constructed by λ -Red mediated homologous recombination using the *cat*
11 or *kan* cassettes from plasmid pKD3 or pKD4, respectively, as the selection marker
12 and primers listed in Table S1. The antibiotic marker was excised using a pCP20-Gm
13 or pCP20-Amp plasmid encoding the FLP recombinase. Complemented mutants were
14 generated by cloning the gene of interest into plasmid pSU2718 between *Bam*HI and
15 *Xba*I cut sites and subsequent transformation.

16 **Growth assays.** Bacterial growth was monitored in U-bottomed 96-well
17 microtitre plates using an automated microplate shaker and reader (FluoStar Optima,
18 BMG Labtech). Each well contained 200 μL of culture supplemented with Cu and/or
19 L-amino acids. Stocks of amino acids were prepared immediately before use. The
20 microplate was sealed with a sterile gas-permeable polyurethane membrane (Sigma).
21 OD_{600} values were recorded up to 16 h. Microplates were shaken at 200 rpm in the
22 orbital mode between readings.

23 **Biochemical analyses.** Batch cultures (50 mL) were prepared in acid-washed
24 glass flasks. Cu was added to the desired final concentration. Bacteria were cultured
25 to the mid-exponential phase ($\text{OD}_{600} \sim 0.3$, ca. 4 doublings) and harvested by

centrifugation (4000 g, 4 °C). The final pellets were processed further for the measurements of intracellular Glu contents, enzyme activities, and gene expression levels. Details are available in the SI Appendix.

Statistical analysis. All statistical analyses were performed using two-way ANOVA in GraphPad Prism7. Results were not corrected for multiple comparisons.

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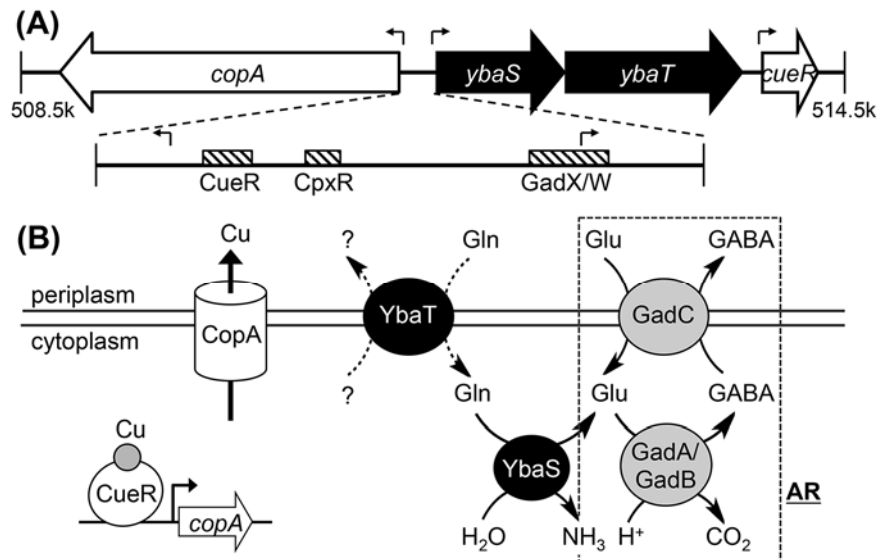
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1 **FIGURES**

2



3

4 **Figure 1. Clustering of Cu tolerance and acid tolerance genes in *E. coli*. (A)**

5 **Genomic context of *copA* and *cueR* in *E. coli*.** Figure shows the approximate

6 locations of genes on the reference genome (RefSeq NC_000913.3). Block arrows

7 represent directions of open reading frames. Transcription start sites are indicated by

8 bent line arrows. Striped boxes represent binding sites for transcription factors (CueR,

9 accttcagcaaggggaaggt; CpxR, gtaaaagtccgtaaa; GadX/W,

10 taaatcaggatgcctgaaaatcggcaccggggtg). **(B) Biochemical function of CopA, CueR,**

11 **YbaS, and YbaT.** CueR is a Cu sensor while CopA is a Cu efflux pump. Both

12 constitute the central mechanism for Cu tolerance in *E. coli*. YbaS is a glutaminase

13 while YbaT is a putative Gln-importing permease. The dashed box shows components

14 of the Glu-dependent acid resistance system (AR), namely the glutamate

15 decarboxylases GadA and GadB, as well as the Glu/GABA-antiporter GadC. Both

16 YbaS and YbaT are thought to support the function of this AR.

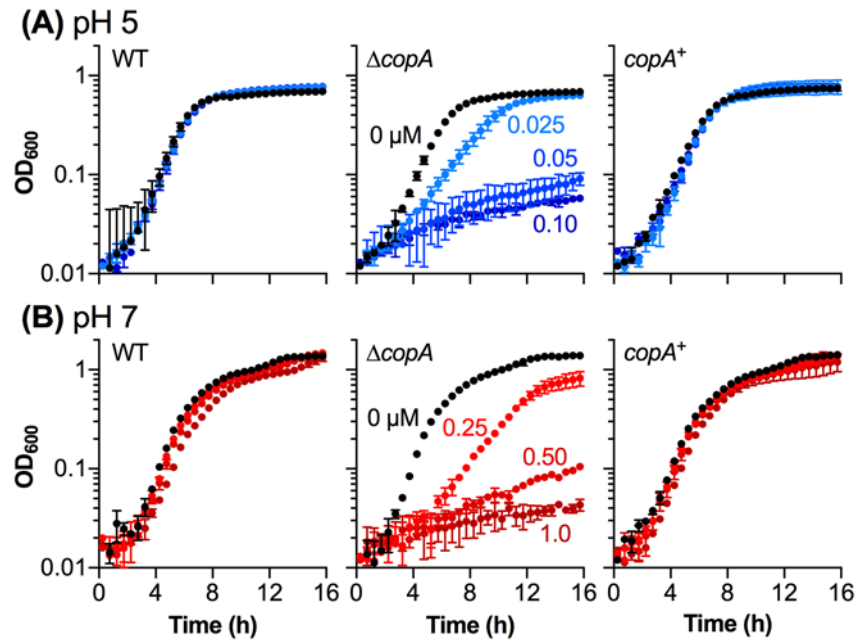


Figure 2. Cu stress during growth under different pH conditions. Growth of *E. coli* EC958 WT, $\Delta copA$ mutant, and $copA^+$ complemented mutant: **(A)** at pH 5 in the presence of 0 – 0.10 μM added Cu and **(B)** at pH 7 in the presence of 0 – 1.0 μM added Cu. Data were averaged from three independent experiments. Error bars represent \pm SD.

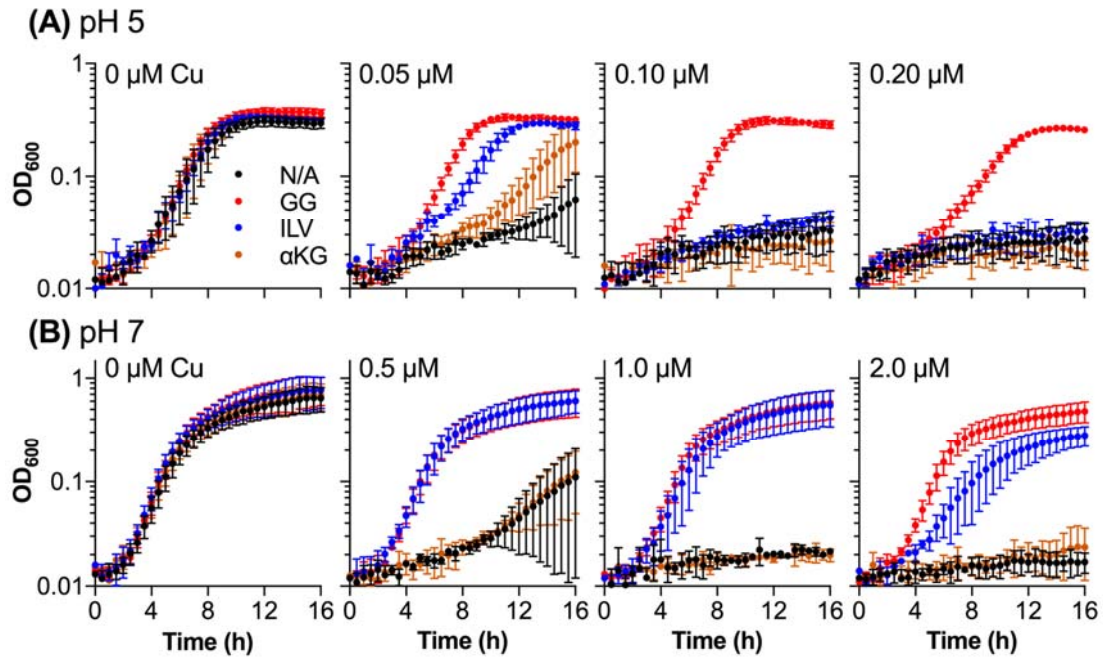


Figure 3. Protective effects of Glu and Gln. *E. coli* UTI89 $\Delta copA$ mutant was cultured (A) at pH 5 in the presence of 0 – 0.2 μM added Cu or (B) at pH 7 in the presence of 0 – 2.0 μM added Cu. The medium was supplemented with water (black, N/A); Glu and Gln (red, GG); Ile, Leu, and Val (blue, ILV); or α -ketoglutarate (orange, αKG). The total concentration of amino acids in each experiment was 0.5 mM. Data were averaged from three independent experiments. Error bars represent \pm SD.

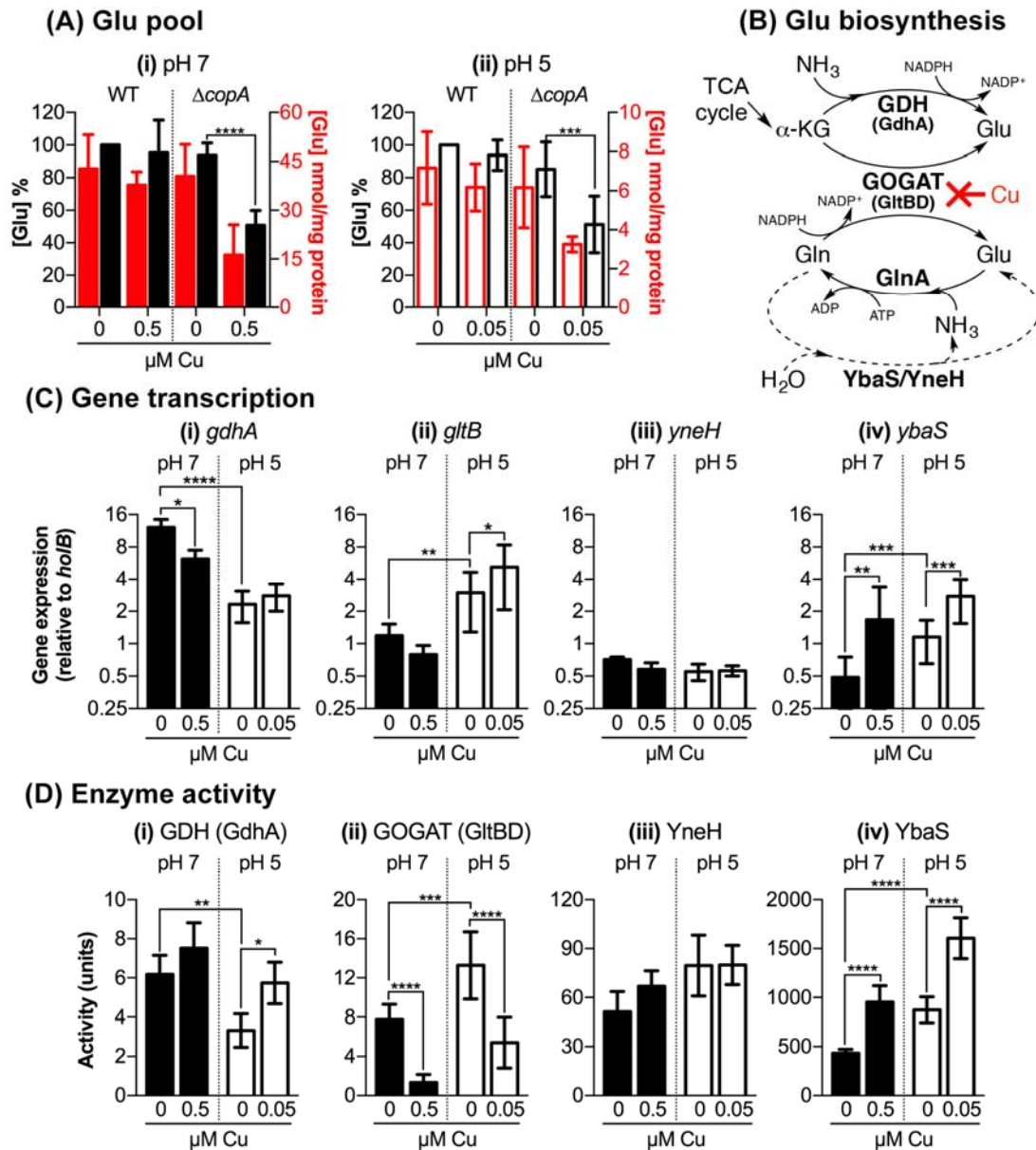


Figure 4. Effects of Cu on glutamate biosynthesis in *E. coli*. (A) Intracellular Glu concentrations. *E. coli* UTI89 WT and $\Delta copA$ mutant were cultured at (i) pH 7 or (ii) pH 5 with or without added Cu as indicated. Intracellular concentrations of Glu were shown as absolute values (red columns) or as a percentage relative to untreated WT (black columns). Data were averaged from five independent biological replicates. (B) Glu biosynthesis pathways. *De novo* synthesis of Glu begins with α -KG from the TCA cycle. This process is catalysed either by GDH using ammonia as the nitrogen donor (top pathway) or by GOGAT using Gln as the nitrogen donor (middle

1 pathway). Glu can also be generated by the hydrolysis of Gln. This process is
 2 catalysed by YbaS or YneH (bottom pathway). Our data show that excess Cu inhibits
 3 biosynthesis at the GOGAT step. The figure also shows one route of Glu consumption
 4 *via* GS, which also generates Gln. **(C) Activities of Glu biosynthesis genes.** The
 5 $\Delta copA$ mutant was cultured at pH 7 (black columns) or pH 5 (white columns) with or
 6 without added Cu as indicated. Activities of **(i)** GDH, **(ii)** GOGAT, **(iii)** YneH, and
 7 **(iv)** YbaS were measured in cell-free lysis extracts. Data were averaged from six
 8 independent biological replicates. **(D) Activities of Glu-synthesizing enzymes.** The
 9 $\Delta copA$ mutant was cultured at pH 7 (black columns) or pH 5 (white columns) with or
 10 without added Cu as indicated. Amounts of **(i)** *gdhA*, **(ii)** *gltB*, **(iii)** *yneH*, and **(iv)**
 11 *ybaS* transcripts relative to *holB* were measured by qPCR. Data were averaged from
 12 six independent biological replicates. **(A), (C), and (D)** Error bars represent \pm SD.
 13 **** $P < 0.0001$, *** $P < 0.001$, ** $P < 0.01$, * $P < 0.05$.

14

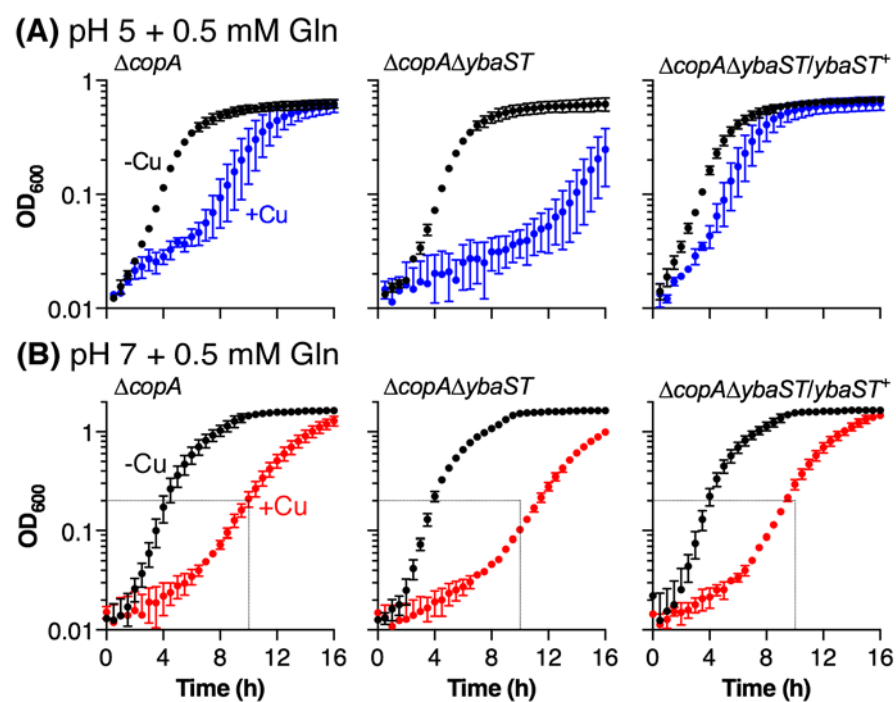


Figure 5. Effects of $\Delta ybaST$ mutation on Gln-dependent Cu tolerance. *E. coli* EC958 $\Delta copA$ mutant, $\Delta copA\Delta ybaST$ double mutant, and $\Delta copA\Delta ybaST/ybaST^+$ complemented mutant were cultured (A) at pH 5 in the presence of 0 (black, -Cu) or 0.05 (blue, +Cu) μM added Cu or (B) at pH 7 in the presence of 0 (black, -Cu) or 0.5 (red, +Cu) μM added Cu. The medium was supplemented with 0.5 mM Gln. The inoculum, used in each experiment was pre-cultured in the same pH. Data were averaged from three independent experiments. Error bars represent \pm SD.

SI APPENDIX

Interplay between tolerance mechanisms to copper and acid stress in *Escherichia coli*

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SI Discussion 1.

Fig. S14 suggested that the amount of Cu required to directly inhibit the activity of mature GOGAT enzyme was higher at pH 5 (10 μ M vs. 1 μ M at pH 7). Earlier, we showed that expression of *gltB* was higher at pH 5 (Fig. 4C(ii)). Similarly, GOGAT activities were higher at pH 5 (Fig. 4D(ii), Figure S10). Therefore, the intracellular concentration of GOGAT enzyme was likely higher in cells cultured at pH 5. Consistently, a higher amount of Cu was required to inhibit GOGAT activity directly at pH 5.

These results seemed to contradict results in Fig. 4D(ii), in which lower amounts of Cu were required to achieve inhibition of GOGAT during growth at pH 5 (0.05 μ M vs. 0.5 μ M at pH 7). However, in these earlier experiments, Cu was added to the culture at the start of the growth and GOGAT activity was measured only when cells reached the mid-exponential phase (after at least 4 generations). The observed loss in GOGAT activity here would be the combined effect of direct enzyme inactivation (as shown in Fig. S14) as well as inhibition of downstream processes that may affect enzyme biogenesis, *eg. via* the Isc pathway for Fe-S cluster assembly (as shown Fig. S12).

SI Discussion 2.

The $\Delta copA \Delta cueR$ mutant was found to be reproducibly more Cu-tolerant at pH 7 compared with the $\Delta copA$ parent strain when growth was assessed in parallel (Fig. S15). The mechanism for the increased tolerance at pH 7 is unknown and is beyond the scope of our work.

At pH 5, the $\Delta copA \Delta cueR$ double mutant was reproducibly more Cu-sensitive than was the $\Delta copA$ parent strain (Fig. S15). On several occasions, growth of the $\Delta copA \Delta cueR$ mutant was affected even in the absence of added Cu and background expression levels of *ybaS* and *ybaT* genes were higher than usual. This was likely a consequence of trace Cu in the

culture medium, which varied between 10–20 nM. These basal amounts of Cu approached inhibitory levels and the precise threshold varied with medium preparations.

To determine if Cu stress induced the expression of *ybaS* and *ybaT* in the $\Delta copA\Delta cueR$ mutant, we cultured this mutant in the presence of 30 nM Cu. Parallel experimentation with the $\Delta copA$ mutant (in the same media preparation, in the presence of 50 nM Cu as usual) as a positive control yielded consistent results and *ybaS* and *ybaT* were reproducibly upregulated during Cu stress (Fig. S16). However, given the challenges described above, it was difficult to obtain reproducible results with the $\Delta copA\Delta cueR$ mutant. On the basis that we were able to find three independent replicates where *ybaS* and *ybaT* were upregulated by Cu (Fig. S16), we concluded that CueR does not control *ybaS* and *ybaT* directly under our experimental conditions.

SI Methods

Bioinformatic analyses. The nucleotide sequences of the *copA-ybaST-cueR* locus from *E. coli* K-12 str. MG1655 (6526 bp, RefSeq NC_000913.3) and the *copA-cueR* locus from *S. enterica* subsp. *enterica* sv. Typhimurium str. LT2 (3029 bp, RefSeq NC_003197.1) were used to query all complete genomes on NCBI (5975 available, last accessed 24/11/2016) using MegaBLAST (v.2.4.0+). Results were visualized using ggplot2 (v.2.2.0) in R (v.3.3.1). Distribution of YbaS was determined by tblastn against the same database with a threshold of 60% identity and 80% coverage. Genomic context was illustrated using Easyfig (1) with manual modification using Inkscape (v.0.91).

Measurement of intracellular Glu content. Bacterial pellets from batch cultures (50 mL) were resuspended in MeOH/MeCN/H₂O (40/40/20 v/v/v %) with frequent vortexing and re-centrifuged. The supernatant was evaporated to dryness under vacuum at 40 °C and the

resulting pellet was resuspended in water. Insoluble debris were removed by centrifugation and the supernatant was added to a reaction mixture containing hydrazine (250 mM), ADP (1 mM), NAD^+ (1.6 mM), and *L*-glutamic dehydrogenase (Sigma G7882, 160 $\mu\text{g/mL}$) in Tris-Cl buffer (100 mM, pH 9). The mixture was incubated at 37 °C for 30–60 min. Glu concentrations in the samples were estimated by comparing final absorbance values at 340 nm against a standard curve. Glu levels in UTI89 strains cultured at pH 7 and pH 5 were routinely measured to be ~44 and ~9 nmol/mg protein, respectively. A parallel culture in Gutnick medium (2) without any added Cu yielded ~90 nmol Glu/mg protein, suggesting that our culture conditions in modified M9 medium were Glu-limiting, particularly at pH 5.

Measurements of enzyme activities. Centrifuged bacterial pellets from batch cultures (50 mL) were resuspended in 0.5 mL of Na-HEPES buffer (50 mM, pH 7.4) and lysed by sonication (5×10 s bursts, 10 W each). Each lysate was centrifuged and the supernatant was added into the appropriate reaction mixture in Na-HEPES buffer (50 mM, pH 7.4) as described below. All reactions were performed at 37 °C. Amounts of proteins in samples were quantified using QuantiPro™ BCA Assay Kit (Sigma).

GOGAT and GDH activities were determined by following the oxidation of NADPH (0.25 mM). Gln (2.5 mM) and α -KG (2.5 mM) were used as substrates for GOGAT. Absorbance values at 340 nm were monitored continuously for 2 min (1 U = 10 nmol NADPH oxidised/min/ μg protein). Gln was replaced with ammonium chloride (25 mM) for GDH (1 U = 1 nmol NADPH oxidised/min/ μg protein). The activity of NUO was estimated by following the oxidation of deamino-NADH (Sigma N6756, 0.25 mM) at 340 nm for 2 min (1 U = 1 nmol deamino-NADH oxidised/min/mg protein). SDH activity was determined by monitoring the reduction of thiazolyl blue tetrazolium bromide (MTT, 0.15 mM) at 570 nm for 15 min in the presence of sodium succinate (5 mM) and phenazine methosulfate (PMS,

0.5 mM) (1 U = 1 nmol of MTT reduced/min/mg protein). To estimate the activities of YbaS and YneH, cell-free lysate supernatants were incubated with Gln (50 mM) at 37 °C for 20 min in MES (100 mM, pH 5.5) and HEPES (100 mM, pH 7.4) buffer, respectively. The mixture was heated to 75 °C for 5 min to deactivate all enzymes and chilled to 4 °C. Enzyme activities were estimated from the amount of Glu generated in the reaction (1 U = 1 nmol Glu hydrolysed/min/mg protein).

RNA extraction and measurement of gene transcripts. For RNA extraction, 1 mL of the batch culture was harvested separately by centrifugation (15,000 g), snap-frozen, and stored at –80 °C until further use. Bacterial RNA was extracted using the RNeasy Mini Kit (QIAGEN) and treated with DNaseI using the RNase-Free DNase Set (QIAGEN). cDNA was generated from 0.5 µg of RNA using the SuperScript® III First-Strand Synthesis System (Invitrogen). qPCR analyses were performed in 10 µL reactions using 2 ng of cDNA as template and 0.4 µM of the appropriate primer pairs (Table S2). Each sample was analysed in three technical replicates. Amplicons were detected with SYBR Green 2 in a QuantStudio 6 Flex Real-Time PCR System (Applied Biosystems). C_q values were calculated using LinRegPCR (3) after correcting for amplicon efficiency. *holB*, which encodes for DNA polymerase III, was used as the reference gene as its expression was not affected by metal ions (4).

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SI TABLES

Table S1. Primers used for making mutant strains. All primers were purchased from Integrated DNA Technologies (Australia). Sequences belonging to the *cat* and *kan* cassettes from pKD3 and pKD4 plasmid, respectively, are in lowercase. Sequences belonging to *E. coli* strain UTI89 (RefSeq NC_007946.1) or EC958 (RefSeq NZ_HG941718.1) are in UPPERCASE. Restriction sites are in **bold**. Ribosomal binding sites are underlined.

Target	Flanking region	Primer Name	Sequence (5' → 3')
	5'	F	CTTTGTCCTGTACCGCCTGC
		R	<u>ggaataggaactaaggagga</u> GGTCAGGTCGATAGTTTGTG
	3'	F	cctacacaatcgctcaagacGTGAGTAACGCCAACCGCTT
		R	AAATGGGTTAATGGCAAGGC
	5'	F	CCTCAGAAACCGCTGTCAGT
		R	<u>ggaataggaactaaggagga</u> TTTGTGTTGCATCTAACATCTTTTGT
	3'	F	cctacacaatcgctcaagacATGAAGCGCAATAAAACCGTATAAC
		R	TTTTCGATAATCGGGCAGTC
	5'	F	AAATGGGTTAATGGCAAGGC
		R	<u>ggaataggaactaaggagga</u> GTGAGTAACGCCAACCGCTT
	3'	F	cctacacaatcgctcaagacATGAAGCGCAATAAAACCGTATAAC
		R	TTTTCGATAATCGGGCAGTC
	5'	F	TCATGGACAACATGGGCAGC
		R	<u>gaagcagctccagcctacaca</u> TCGCTAATGTTTCATCGTTTCG
	3'	F	ctaaggaggatattcatatgCTCTCCGGCTGCTGTCAT
		R	ACAGCGTCAGACGGCTATTT
pSU2718 - <i>copA</i>	-	F	<i>catgga</i> GGATCC <u>ttaaagaggagaaaggtaccgc</u> ATGTCACAAACTATCGACCTGAC
		R	<i>ttacgc</i> TCTAGACGC ATCCGCAATGATGTACT
pSU2718 - <i>ybaST</i>	-	F	<i>catgga</i> GGATCC <u>ttaaagaggagaaaggtaccgc</u> ATGTTAGATGCAAACAAATTACAGC
		R	<i>ttacg</i> CTCTAGATGTGCTTTGT TAAAGGGTTTCA

^aThese primers were used to generate the $\Delta copA$ mutant in both UTI89 and EC958 backgrounds.

^bThis mutation was introduced in the UTI89 $\Delta copA$ background to generate UTI89 $\Delta copA\Delta cueR$ double mutant.

Table S2. Primers used for analyses of gene transcription by qPCR. Primers were purchased from Sigma (Australia). Genome sequence of *E. coli* strain UTI89 (RefSeq NC_007946.1) was used as template.

Amplicon name	Primer sequence (5' → 3')	Amplicon length (bp)
<i>copA</i>	F: ATGACGATGACAGCCAGCAG R: TTTACCCGTGCCTGAGTGAC	112
<i>cueO</i>	F: AACCAGTGAAGGTGAGCGAG R: CATCCCCATCTGACTGACCG	119
<i>cueR</i>	F: GAAGAGAAGGGGCTGGTGAC R: CCTCCAGGTAAAGCCCACC	124
<i>gltB</i>	F: AAGGTCTGTGTATGCCGACG R: GCGGTACGGTGTTAGTGGAG	114
<i>holB</i>	F: GTGGTGCGAAAGTTGTCTGG R: CGCGGGTAGCAAGGAAAAAC	120
<i>sufA</i>	F: GCTTTGGCTATGTGCTCGAC R: CTTGCAGCGGGACAAACAG	93
<i>sufB</i>	F: TGGAGATGGAAGAACCGCAC R: GTCACAATTACCGCACGATGG	98
<i>ybaS</i>	F: ACCAACTTCCATAACCGGGC R: TTGATGAGCGTGGAGCACTG	113
<i>ybaT</i>	F: TAAGCAACCATGCGGTAGGG R: AGAAACGGAAATATGCGCCG	119

SUPPORTING FIGURES

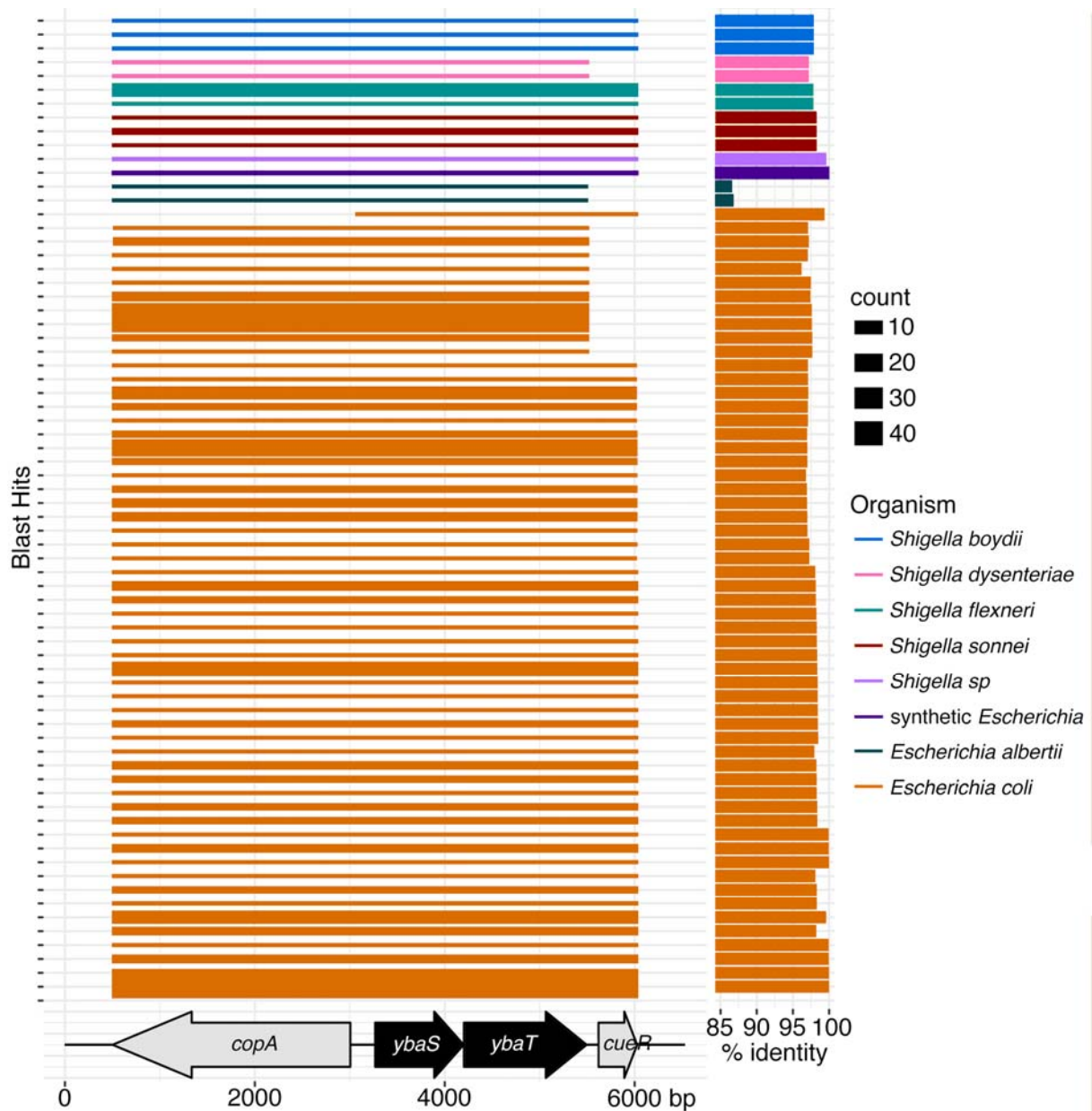


Figure S1. Sequence alignments of *copA*-*ybaST*-*cueR* loci. The sequence of the *copA*-*ybaST*-*cueR* locus from *E. coli* was used as a query in a BLASTn search against 5975 complete bacterial genomes. The search yielded a total of 232 positive hits with alignment length covering *ybaST*. Each horizontal line represents a unique alignment group belonging to the same species that has the same % identity over the same alignment length. The line thickness represents the number of hits in each group and is coloured according to species name.

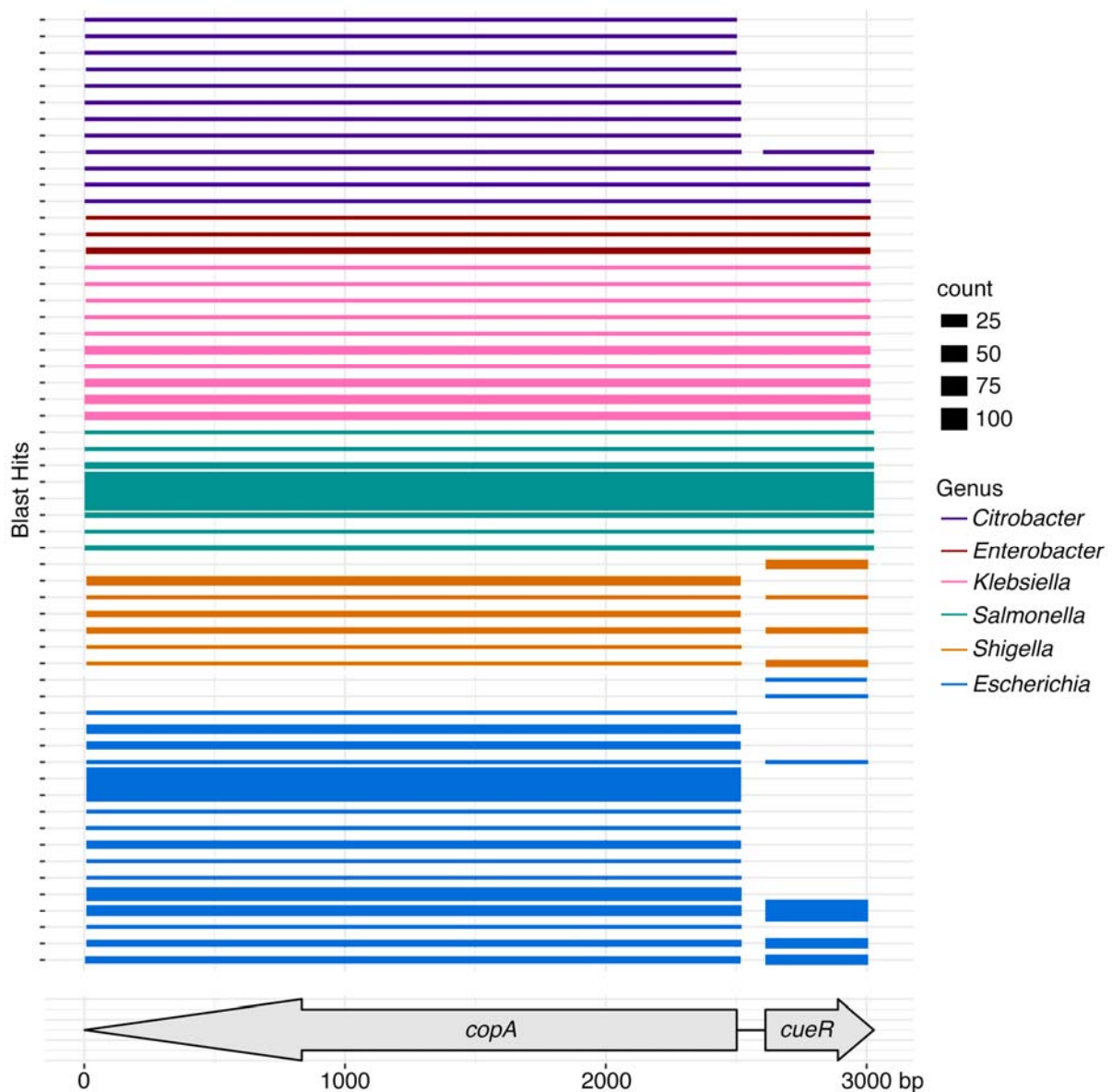


Figure S2. Sequence alignments of *copA-cueR* loci. The sequence of the *copA-cueR* locus from *S. enterica* sv. Typhimurium was used as a query in a BLASTn search against 5975 complete bacterial genomes. The search yielded a total of 262 positive hits with alignment length covering *copA* and *cueR* without *ybaST* insertion, shown by continuous horizontal lines with no gap. Hits from *Escherichia* and *Shigella* genera, which include *ybaST* insertion were also shown by horizontal lines with a gap between *copA* and *cueR* for comparison. Each horizontal line represents a unique alignment group belonging to the same species that has the same % identity over the same alignment length. The line thickness represents the number of hits in each group and is coloured according to species name.

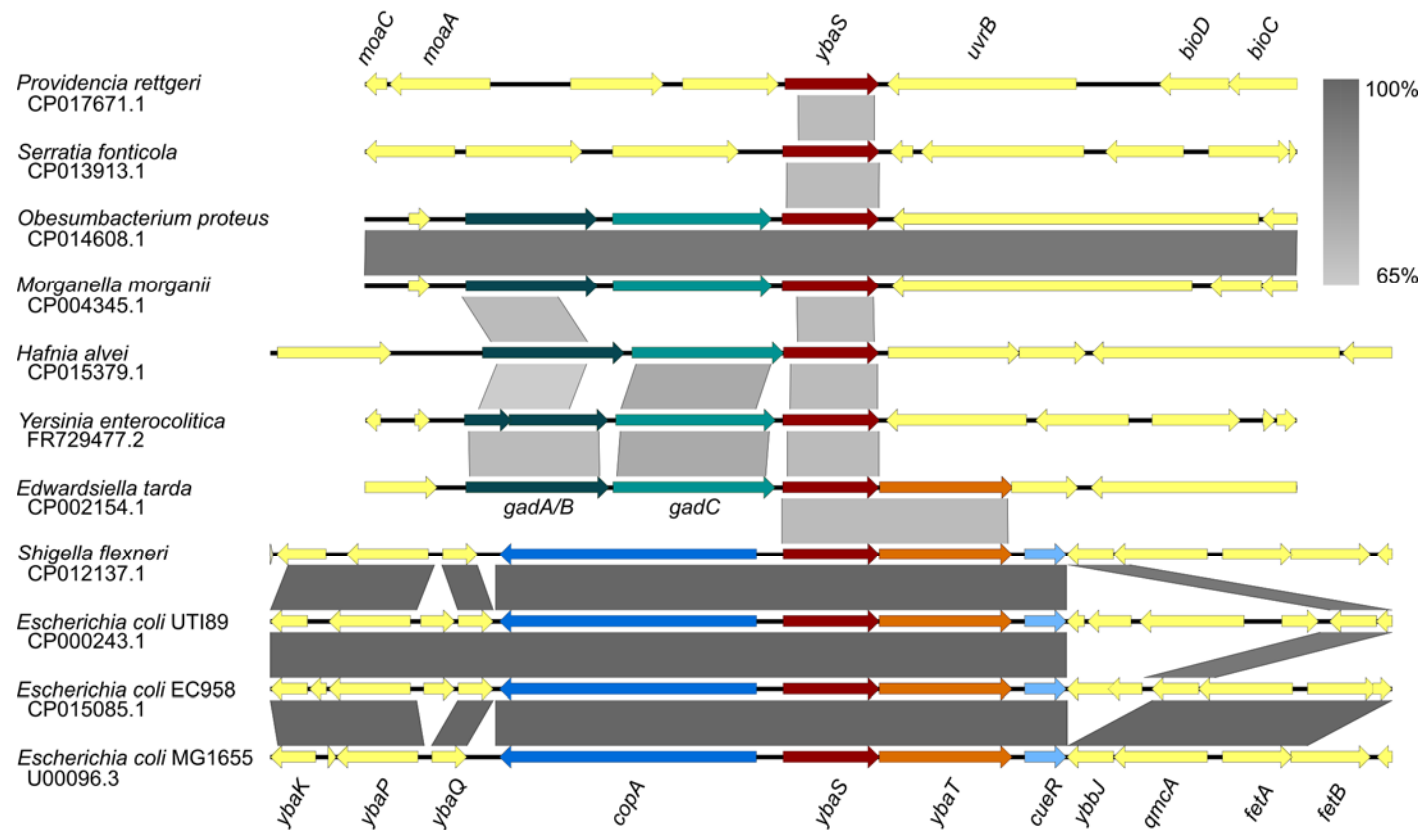


Figure S3. Genomic context of *ybaS* in several *Enterobacteriaceae* species. The protein sequence of YbaS was used to query 5975 complete bacterial genomes by tblastn. The search identified 325 genomes positive for YbaS (>60% identity over 80% coverage) from 15 genera. One genome per genus from *Enterobacteriaceae* family was chosen to represent the YbaS-encoding locus plus 5 kb flanking regions. The *copA-ybaS-ybaT-cueR* arrangement is confined within *E. coli* and *Shigella* while *ybaS* is located elsewhere in the genomes of other genera.

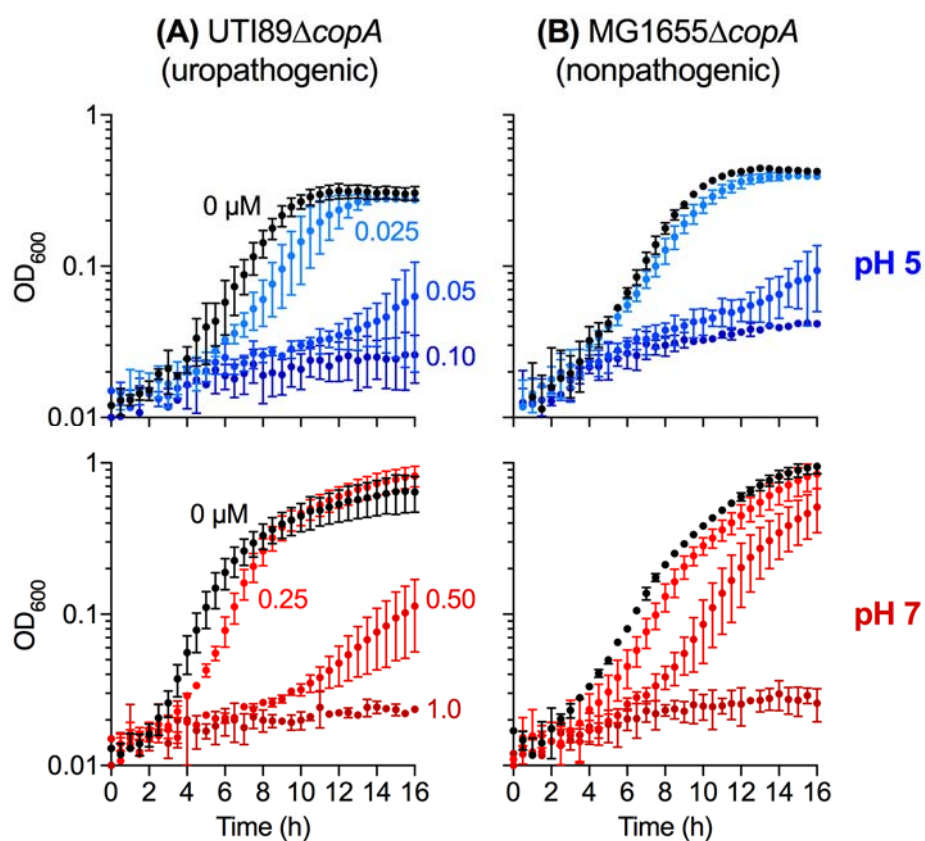


Figure S4. Cu stress in different *E. coli* strains. The $\Delta copA$ mutant strains of (A) UTI89 and (B) K-12 substr. MG1655 were cultured at pH 5 and pH 7 in the presence of various concentrations of Cu as indicated. Data were averaged from three independent experiments. Error bars represent \pm SD.

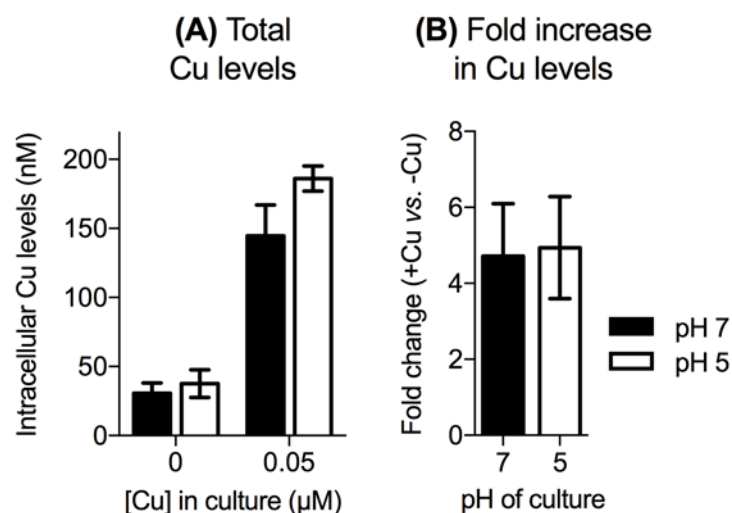


Figure S5. (A) Total amounts of intracellular Cu in *UTI89ΔcopA*. Bacteria were cultured at pH 7 (black columns) or pH 5 (white columns) to a final OD₆₀₀ of ~0.3. The medium was supplemented with 0 or 0.05 μM Cu as indicated. Bacteria were centrifuged, washed once with ice-cold PBS containing 1 mM EDTA and twice more with ice-cold PBS without any EDTA. Bacterial pellets were dried overnight at 50 °C and dissolved in conc. nitric acid (100 μL) with heating at 90 °C. Each sample was diluted to 10 mL with deionised water and sent for metal analyses using Inductively-Coupled Plasma Mass Spectrometry at the School of Earth Sciences, The University of Queensland. Cu levels were normalised to protein content. Intracellular Cu concentrations (in nM) were calculated using the assumption that the protein concentration inside an *E. coli* cell is ~300 mg/mL. **(B) Fold increase in intracellular Cu levels.** This was calculated using data from panel (A). **(A-B)** Results were averaged from five independent experiments. Error bars represent ± SD.

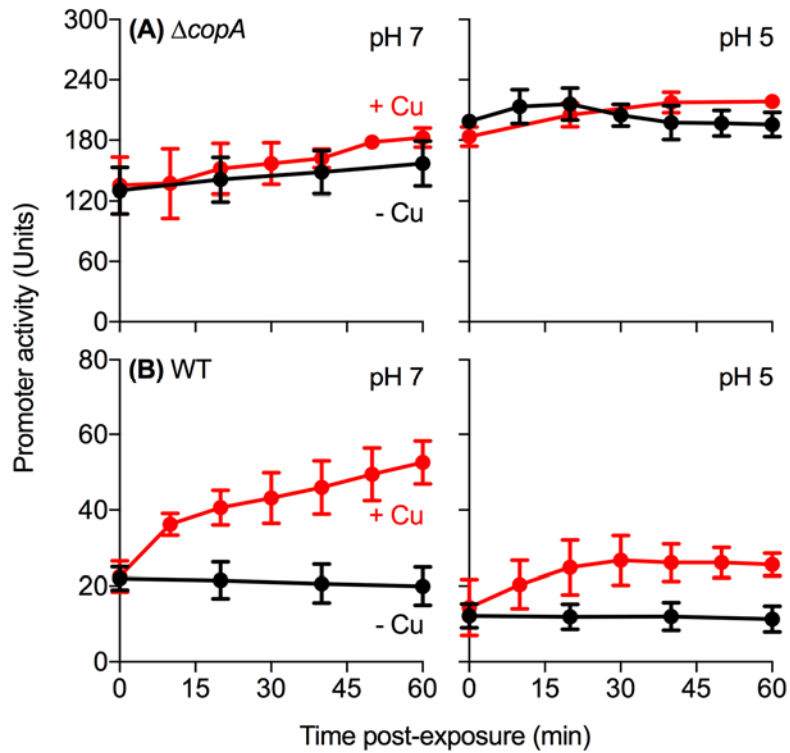


Figure S6. Response of $P_{copA-lacZ}$ reporter to Cu stress at pH 7 and pH 5. *E. coli* UTI89 (A) $\Delta copA$ mutant and (B) WT strains harbouring the $P_{copA-lacZ}$ plasmid were cultured at pH 7 or pH 5 without any added Cu. Upon reaching the mid-exponential phase, bacteria were challenged with water (-Cu, black traces) or 1 μ M of added Cu (+Cu, red traces) in the same medium. Bacteria were collected at intervals up to 60 min post-exposure. β -galactosidase activities were measured following standard protocol using *o*-nitrophenyl- β -galactoside (1 mg/mL) as substrate. The total volume of each sample was 200 μ L. Absorbance values at 420 nm were recorded in microtitre plates and results were expressed as Miller units. Data were averaged from three independent experiments. Error bars represent \pm SD.

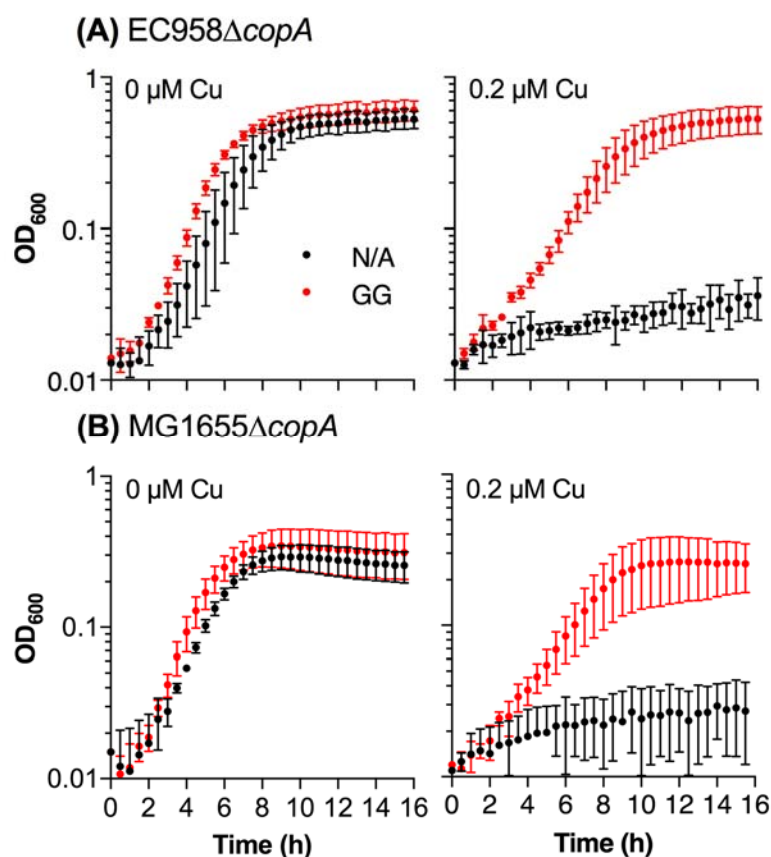


Figure S7. Protective effects of Glu and Gln in different *E. coli* strains. *E. coli* (A) EC958 Δ *copA* and (B) MG1655 Δ *copA* mutant strains were cultured at pH 5 in the presence of 0 or 0.2 μ M added Cu. The medium was supplemented with water (black, N/A) or a combination of Glu and Gln (0.25 mM each, red, GG). Data were averaged from three independent experiments. Error bars represent \pm SD.

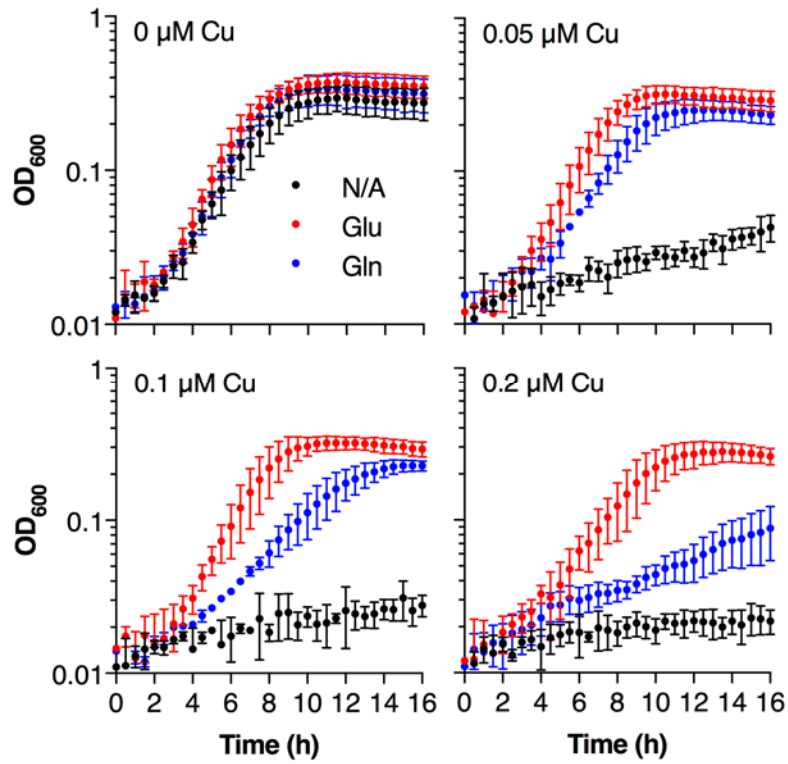


Figure S8. Separate protective effects of Glu and Gln. *E. coli* UTI89Δ*copA* mutant strain was grown at pH 5 in the presence of 0 – 0.2 μM added Cu. The culture medium was supplemented with water (black), 0.5 mM Glu (red), or 0.5 mM Gln (blue). Data were averaged from three independent experiments. Error bars represent ± SD.

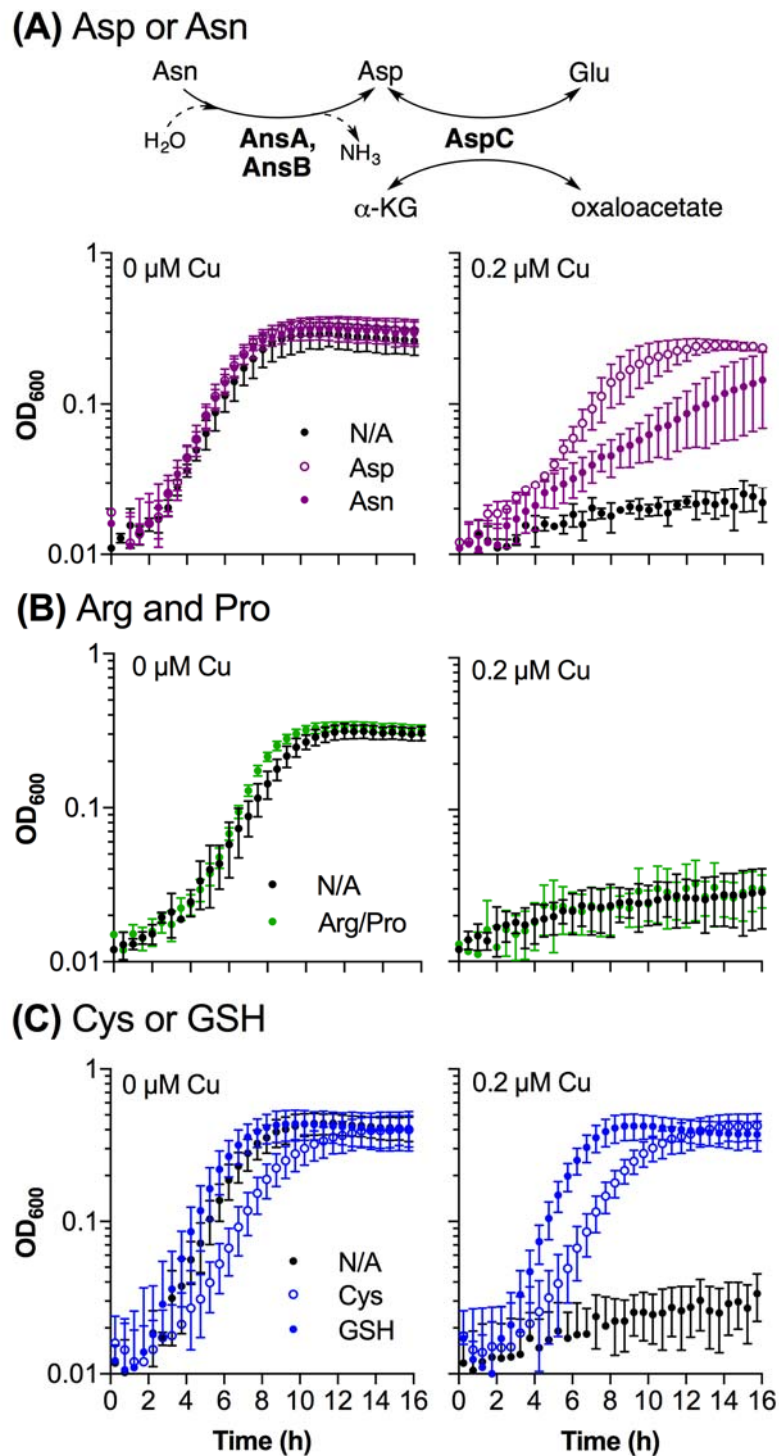


Figure S9. Protective effects of (A) Asp or Asn, (B) Arg and Pro, (C) Cys or GSH. *E. coli* UTI89 Δ *copA* mutant strain was cultured at pH 5 in the presence of 0 or 0.2 μ M added Cu. The medium was supplemented with water (N/A, black circles) or 0.5 mM of each amino acid as indicated. Data were averaged from three independent experiments. Error bars represent \pm SD. Pathway for the generation of Glu from Asn and Asp via the two asparaginases in *E. coli* (AnsA and AnsB) and aspartate aminotransferase (AspC) was shown in panel (A).

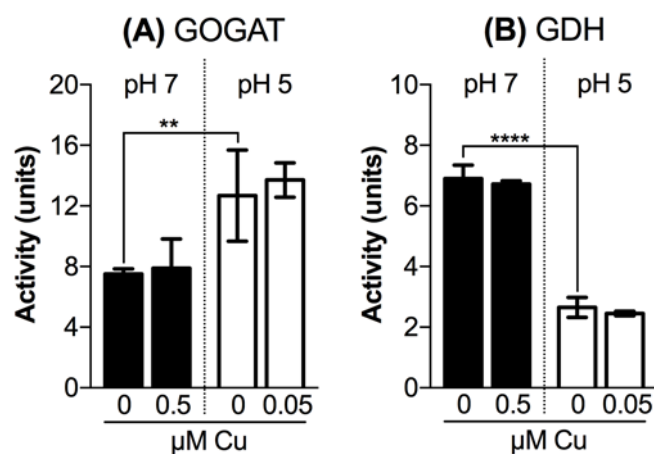


Figure S10. Effects of Cu on the activities of glutamate-synthesising enzymes in WT bacteria. UTI89 WT was cultured at pH 7 (black columns) in the presence of 0 or 0.5 μM Cu, or at pH 5 (white columns) in the presence of 0 or 0.05 μM Cu. Mid-exponential cells were collected and activities of **(A)** GOGAT and **(B)** GDH were measured in cell-free lysis extracts. Data were averaged from three independent experiments. Error bars represent \pm SD. GOGAT activities were reproducibly higher in bacteria cultured at pH 5 (** $P < 0.01$) while GDH activities were higher in bacteria cultured at pH 7 (**** $P < 0.0001$).

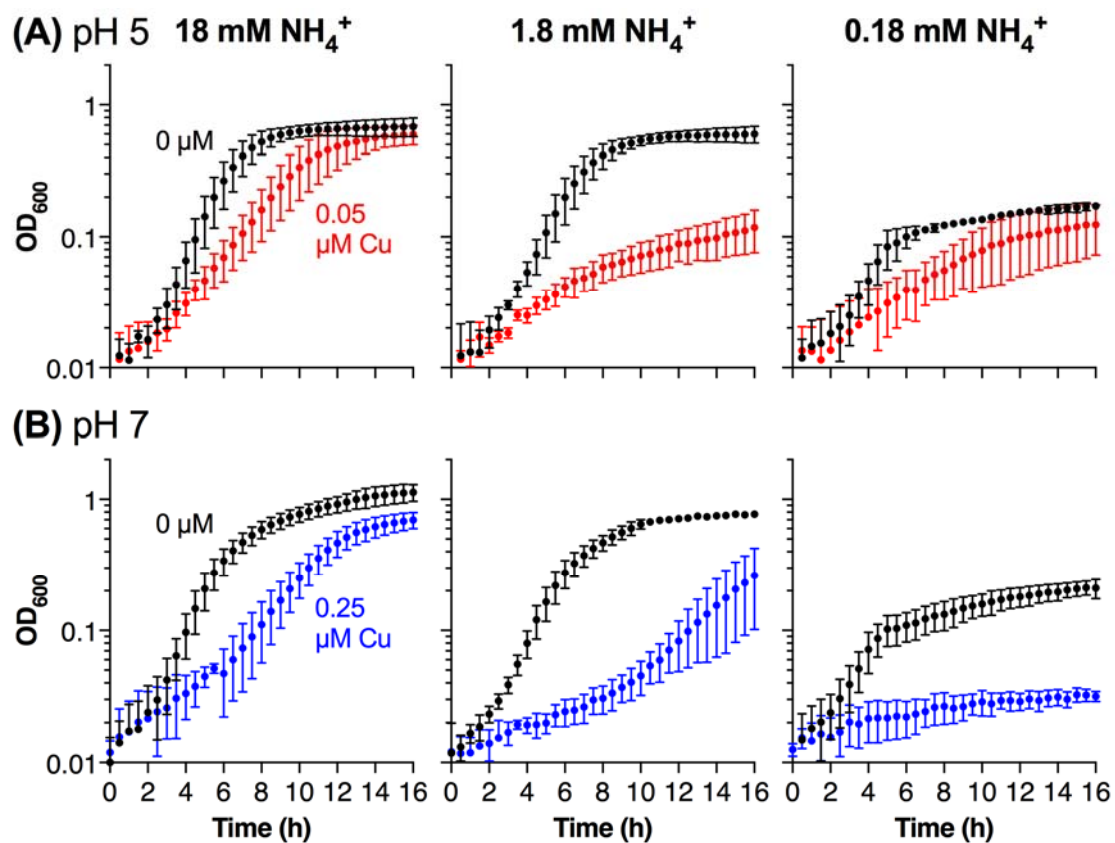


Figure S11. Effects of ammonia availability on Cu stress. *E. coli* UTI89 ΔcopA mutant strain was grown at (A) pH 5 or (B) pH 7 with or without Cu as indicated. The culture medium contained 18, 1.8, or 0.18 mM ammonium chloride as the sole nitrogen source. Data were averaged from four independent experiments. Error bars represent \pm SD.

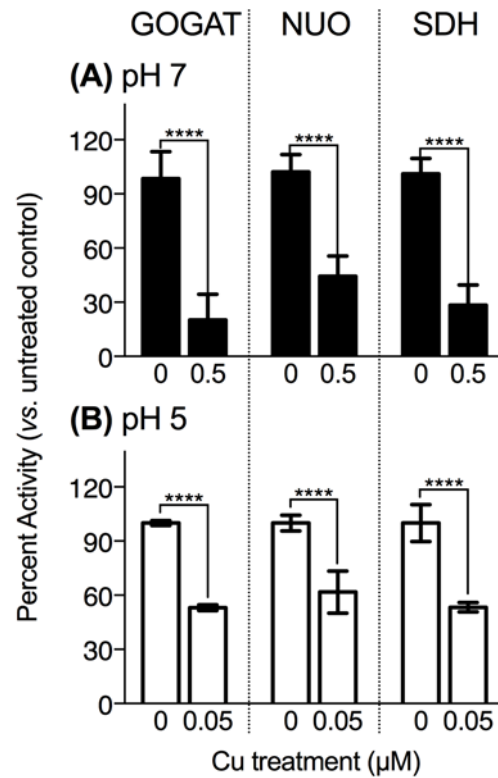


Figure S12. Effects of Cu on the activities of iron-sulfur enzymes. *E. coli* UTI89 Δ *copA* mutant strain was cultured (A) at pH 7 in the presence of 0 or 0.5 μ M added Cu or (B) at pH 5 in the presence of 0 or 0.05 μ M added Cu. Mid-exponential cells were collected and activities of GOGAT, NUO, and SDH were measured in cell-free lysis extracts. Results were normalised to the untreated control (cultured with 0 μ M added Cu). Data were averaged from three independent experiments. Error bars represent \pm SD. **** $P < 0.0001$.

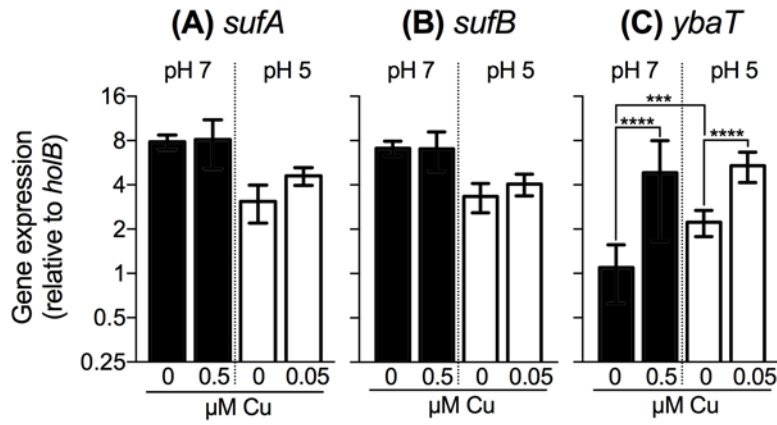


Figure S13. Effects of Cu on expression of (A) *sufA*, (B) *sufB*, and (C) *ybaT*. *E. coli* UTI89 Δ *copA* mutant strain was cultured at pH 7 (black columns) or in the presence of 0 or 0.5 μM Cu at pH 5 (white columns) in the presence of 0 or 0.05 μM Cu. Total RNA was extracted from mid-exponential cells and amounts of transcripts relative to *holB* were measured by qPCR. Data were averaged from six independent experiments. Error bars represent \pm SD from the mean. *** $P < 0.001$, **** $P < 0.0001$.

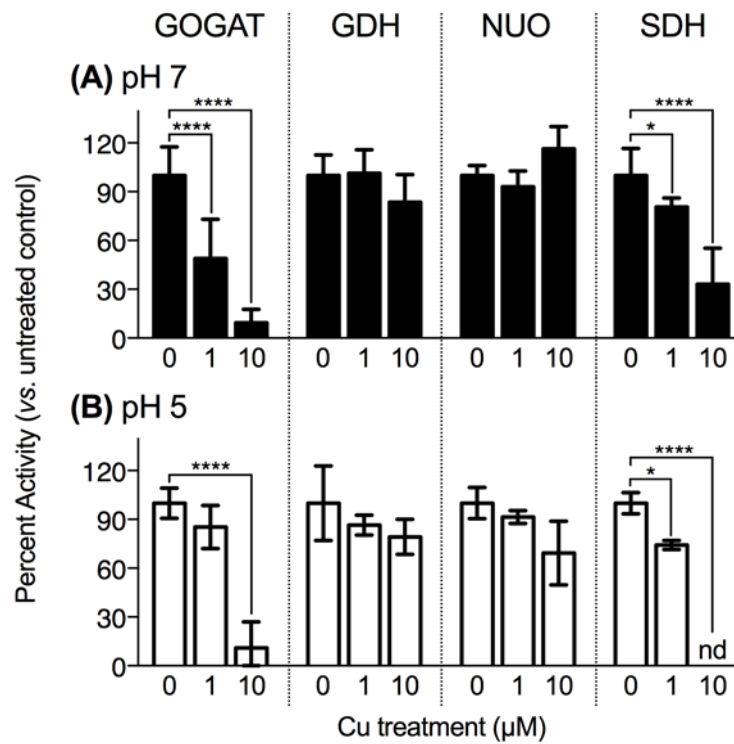


Figure S14. Excess Cu ions may directly damage GOGAT. *E. coli* UTI89 Δ *copA* mutant strain was cultured at (A) pH 7 or (B) pH 5 without any added Cu to the mid-exponential phase and was subsequently challenged with 0, 1, or 10 μ M Cu. After 30 min, cells were collected and activities of GOGAT, GDH, NUO, and SDH were measured in cell-free lysis extracts. Data were averaged from three independent experiments. Error bars represent \pm SD. **** $P < 0.0001$, * $P < 0.05$.

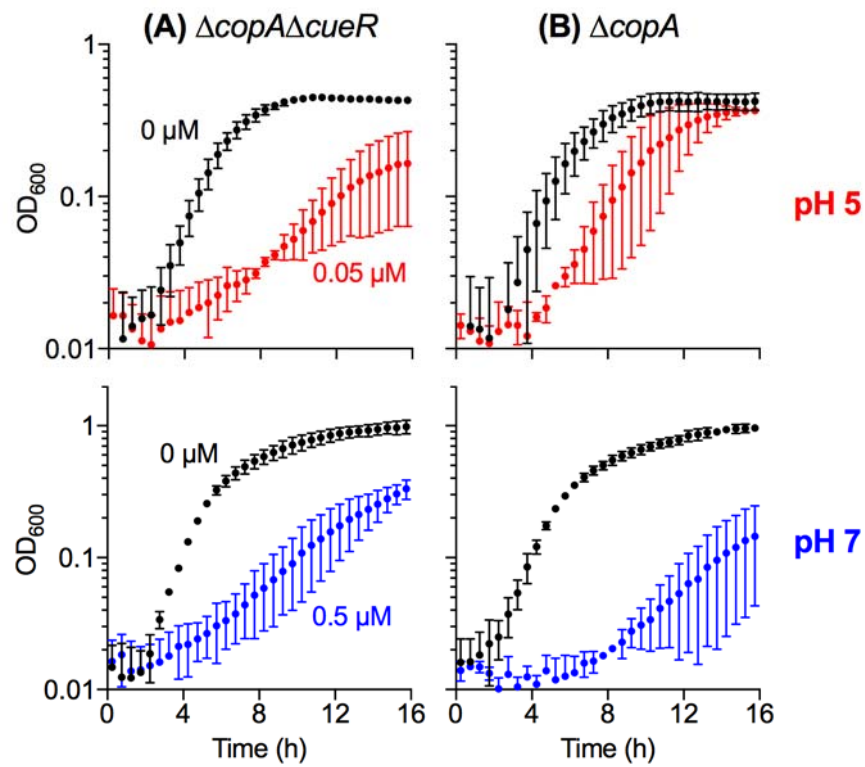


Figure S15. Cu stress in the UTI89Δ*copA*Δ*cueR* mutant. Bacteria were cultured at pH 5 in the presence of 0 or 0.05 μM of added Cu (top panels) or at pH 7 in the presence of 0 or 0.5 μM of added Cu (bottom panels). Data were averaged from three independent experiments. Error bars represent ± SD. Cu stress in the UTI89Δ*copA* parent mutant was assessed in parallel and the results were shown for comparison.

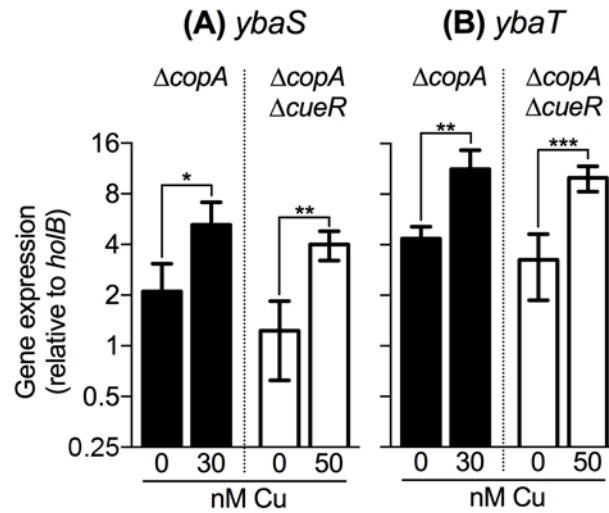


Figure S16. Effects of Cu on expression of (A) *ybaS* and (B) *ybaT* in the UTI89 $\Delta copA \Delta cueR$ mutant. Bacteria were cultured at pH 5 in the presence of 0, 30, or 50 nM of added Cu as indicated. Data were averaged from three independent experiments. Error bars represent \pm SD. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$. Results from parallel experimentation with the UTI89 $\Delta copA$ parent mutant strain were shown for comparison.

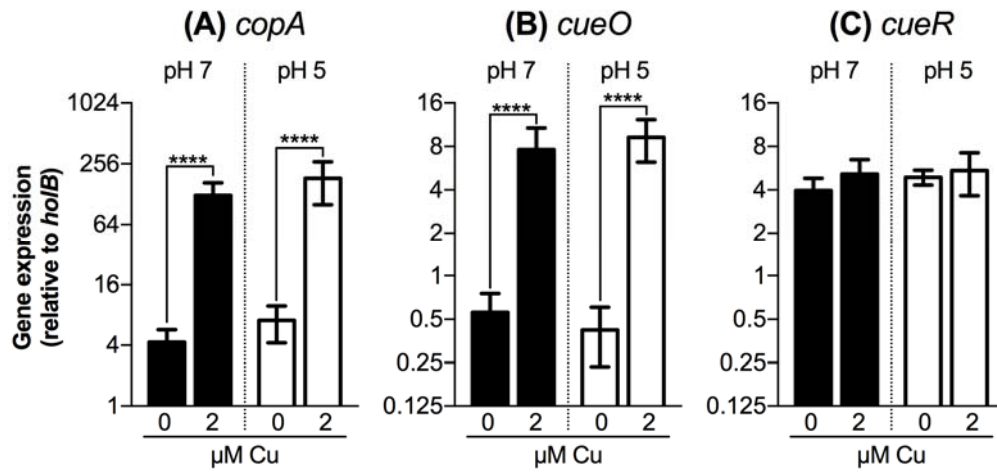


Figure S17. Effects of pH on the expression of Cu tolerance genes in WT cells. The UTI89 WT strain was cultured at pH 7 (black columns) or pH 5 (white columns) in the presence of 0 or 2 μ M Cu. Total RNA was extracted from mid-exponential cells and amounts of (A) *copA*, (B) *cueO*, and (C) *cueR* transcripts relative to *holB* were measured by qPCR. Data were averaged from six independent experiments. Error bars represent \pm SD from the mean. **** $P < 0.0001$.

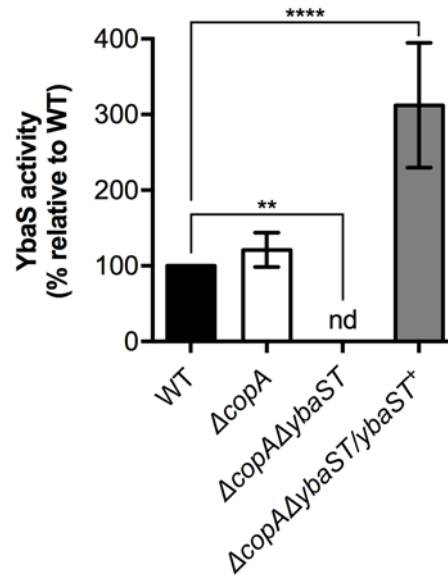


Figure S18. YbaS activity in $\Delta ybaST$ mutant strains. Bacteria were propagated on Cu-free LB agar overnight and cells from the agar plate were harvested and lysed. YbaS activities were measured in cell-free lysis extracts. Results were normalised to the WT. Data were averaged from three independent experiments. Error bars represent \pm SD. nd, not detectable (below detection limit). **** $P < 0.0001$, ** $P < 0.01$.

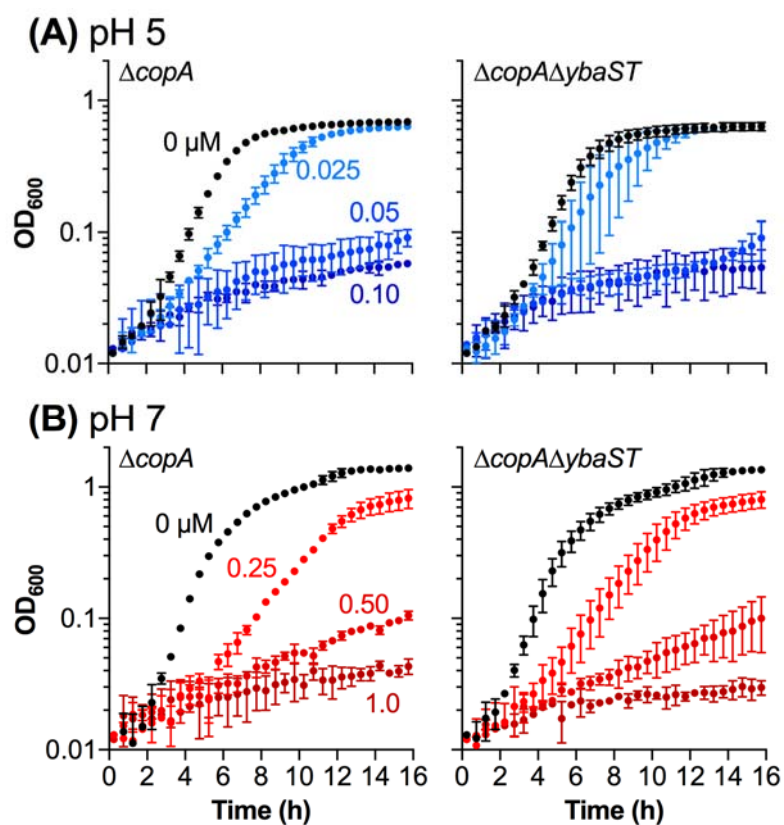


Figure S19. Cu stress in the $\Delta copA\Delta ybaST$ mutant strain. Growth of EC958 $\Delta copA$ and $\Delta copA\Delta ybaST$ mutant strains (A) at pH 5 in the presence of 0 – 0.10 μM of added Cu and (B) at pH 7 in the presence of 0 – 1.0 μM of added Cu. Data were averaged from three independent experiments. Error bars represent \pm SD.

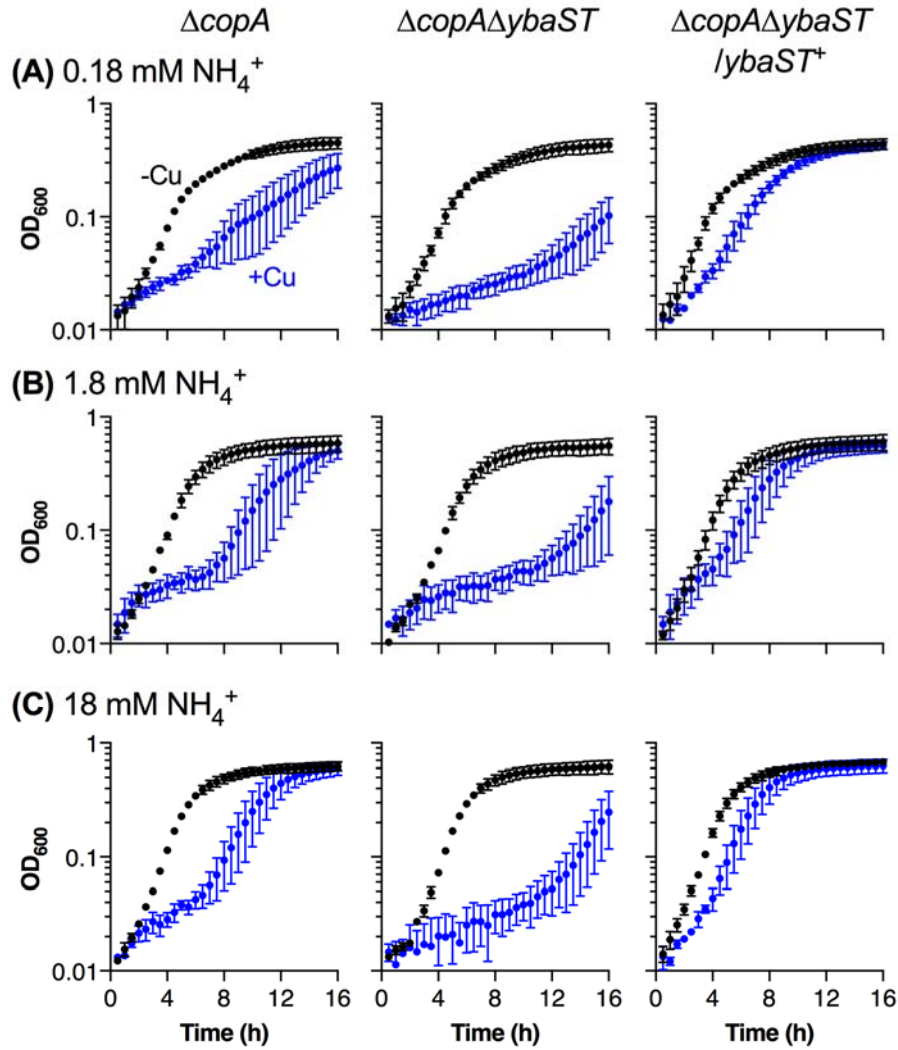


Figure S20. Effects of ammonia availability on Cu stress in the $\Delta copA\Delta ybaST$ mutant. *E. coli* EC958 $\Delta copA$ mutant, $\Delta copA\Delta ybaST$ double mutant, and $\Delta copA\Delta ybaST/ybaST^+$ complemented mutant strain was cultured at pH 5 with 0 (-Cu, black traces) or 0.05 μM of added Cu (+Cu, blue traces). The culture medium contained 0.5 mM Gln and (A) 0.18, (B) 1.8, or (C) 18 mM ammonium chloride. Data were averaged from three independent experiments. Error bars represent \pm SD.